DB 9; Length 1743;

Length 1743;

DB 9;

Length 1743;

DB 9;

Query Match

Query Match

CDNA #226

Length 1743

Length 1743;

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ACF30523 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003067478-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003054458-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ
US2003059886-A1.
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                                                                                 ACD91212 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
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Pred. No. 41;
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Pred. No. 41;
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                                5.4%; Score 38.4; 1
52.5%; Pred. No. 41;
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Pred. No. 41;
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RESULT 688
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(GETH ) GENENTECH INC.
13-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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                                             Best Local Similarity RESULT 684
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Best Local Similarity
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Best Local Similarity
RESULT 691
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SULT 689
                                                                                                                                                                                      Best Local Similarity
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                                                                                                                     US2003049751-A1.
13-MAR-2003.
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                                  Query Match
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054480-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068711-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003049770-A1.
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US2003049745-A1.
 RESULT 674

ID ACD10523 standard, cDNA, 1743 BP.

ID ACD10523 standard, cDNA, 1743 BP.

E Human secreted/transmembrane protein (PRO) cDNA #226.

PN US2003036164-A1.

PD 20-FEB-2003.
                                                                                                                                     ACD12165 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
US2003040074-A1.
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13-MAR-2003.
(GETH ) GENENTECH INC.
6rv Match 5.4%; Score 38.4; D
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
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27-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 680
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RESULT 682
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RESULT 677
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Best Local Similarity
RESULT 675
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RESULT 676
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Query Match

Query Match

9789

Query Match

ID NO:451

Length 1743

Length 1743;

Length 1743;

DB 9;

Length 1743;

DB 9;

Length 1743

DB 9;

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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
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(GETH ) GENENTECH INC.
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Query Match
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(GETH ) GENENTECH INC.
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A (GETH ) GENENTECH INC.
Query Match
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Best Local Similarity
RESULT 707
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Best Local Similarity
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RESULT 708
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Best Local Similarity
RESULT 704
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                 US2003082715-A1
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                                                                          RESULT 703
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 ACF08097 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003049758-A1.
                                                                                                   ACF08404 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003064448-A1.
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                                                                          Length 1743;
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003068756-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003068735-A1.
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                                                                        52.5%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
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Best Local Similarity 52.5%; Pred. No. 41;
RESULT 700
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 698
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RESULT 702
ID ACF52659 Ctradent
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Best Local Similarity
RESULT 695
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Best Local Similarity
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Best Local Similarity
RESULT 701
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003103407-A1. 05-UNA-2003 (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ACF61811 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003100061-A1.
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                                              ACF52966 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003082716-A1.
                                                                                                                                                             ACF64959 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068737-A1.
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                                                                                                             Length 1743;
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Length 1743;
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Human secreted/transmembrane protein (PRO) cDNA #226.
13-PEB-2003.
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US2003034477-A1.
20-MAR-2003.
(GETH ) GENENTECH INC.
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                                                                                                                DB 9;
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Score 38.4; I
Pred. No. 41;
                                                                                                               Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Best Local Similarity 52.5%;
5.4%;
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Query Match

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ACF03282 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451.
US200349744-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
17-APR-2003
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003049783-A1.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104557-A1.
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US2003044920-A1.
06-MAR-2003.
                                                                                                                            ADAB1722 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
US2003092121-A1.
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US2003027276-A1.
06-FEB-2003.
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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US2003027265-A1.
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Best Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 727
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RESULT 726
                                                                                       Best Local Similarity RESULT 722
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ACD22315 standard;
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                                           05-JUN-200
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                                           ACF07483 standard; cDNA; 1743 BP.

Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003049753-A1.
13-MAR-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                              Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003049763-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003073172-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068736-A1.
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Best Local Similarity 52.5%; Pred. No. 41;
RESULT 721
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003068700-A1.
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                                                                                                                                                                                                                                                                                                                             ACF20948 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, US2003073772-A1.
17-APR-2003.
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                                                                                                                                 52.5%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
            52.5%; Pred. No. 41;
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cDNA encoding human PRO polypeptide #226.
US2003073182-A1.
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                                                                                                                                                                                        CDNA; 1743 BP
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 720
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Best Local Similarity
RESULT 714
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Best Local Similarity
       Best Local Similarity RESULT 712
                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 715
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RESULT 716
                                                                                                                                                   Local Similarity
                                                                                                                                                                                        ACF20641 standard;
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Query Match

Best Loca RESULT 719

Query Match

Query Match

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Query Match
Best Local Similarity
RESULT 741
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Best Local Similarity
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RESULT 748
                                                                                                                                                                                                                                  ACH07674 standard;
                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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                                ACF50817 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003032121-A1.
                                                                                                                                                        Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451. US2003064458-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068702-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068734-A1.
                                                                                                                                                                                                                                                                                                                                         Length 1743;
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                                                                                               Length 1743;
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                                                                                                                                                                                                                                                                                                                                                                                ACD48379 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
US2003064464-A1.
                                                                                                                                                                                                                                                         ACD46537 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
US2003064460-A1.
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                                                                                            5.4%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
                                                                                                                                                                                                                  52.5%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                        DB 9;
                                                                                                                                                                                                                                                                                                                                        Score 38.4; | Pred. No. 41;
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52.5%; Pred. No. 41;
    52.5%; Pred. No. 41;
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Human PRO polynucleotide #226.
US2003068738-A1.
                                                                                                                                        ACF34312 standard; cDNA; 1743 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACF27760 standard; cDNA; 1743 BP
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity 5
RESULT 733
ID ACD46537 standar
DE Human secret
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(GETH ) GENENTECH INC.
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RESULT 739
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Best Local Similarity RESULT 731
                                                                                            Query Match
Best Local Similarity
RESULT 732
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Best Local Similarity
RESULT 738
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ACF49282 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104540-Al.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003040059-A1.
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                                                                          Length 1743;
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                                                                                                             Achuvis67 standard; CDNA; 1743 BP.

Human secreted/transmembrane protein (PRO) cDNA #226.
US203049742-A1.
                                                                                                                                                                                                                                                                                             Human secreted/transmembrane protein (PRO) cDNA #226 US2003049747-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003049779-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003040078-A1.
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                                                                        Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Best Local Similarity 52.5%; Pred. No. 41;
RESULT 746
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Pred. No. 41;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ACH11479 standard; cDNA; 1743 BP.
cDNA encoding human PRO polypeptide #226.
US2003049766-A1.
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cDNA encoding human PRO polypeptide #226.
US2003049767-A1.
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                                                                        5.4%;
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(GETH ) GENENTECH INC.
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Best Local Similarity
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                                                                                                                    RESULT 760
ID ACF76297 standard;
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                                                                                                     Best
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Human secreted polypeptide PRO1337-encoding CDNA, SEQ ID NO:451.
US2003073174-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ACF25246 standard; CDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US200368712-A1.
                                                                                                                                                                                    ACF17957 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
                                                                                                                                                                                                                                                                                                                            ACF32740 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003064445-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ACF40401 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003064449-A1.
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Pred. No. 41;
              Length 1743;
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                                                                                                                                                                                                                                                                             5.4%; Score 38.4; DB 9; Length 1743; 52.5%; Pred. No. 41;
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            Ouery Match
S.4%; Score 38.4; DB 9; Leng
Best Local Similarity 52.5%; Pred. No. 41;
RESULT 70.1
ID ACD31456 standard; CDNA; 1743 BP.
DE Human secreted/transmembrane protein (PRO) CDNA #226.
PN US200332132-Al.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003064441-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003068696-A1.
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Pred. No. 41;
                                                                                                                                    52.5%; Score 38.4; I 52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%;
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(GETH ) GENENTECH INC.
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RESULT 753
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Best Local Similarity
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RESULT 755
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RESULT 756
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Best Local Similarity
RESULT 757
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Best Local Similarity
RESULT 759
06-MAR-2003.
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RESULT 754

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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104541-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003032133-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
92200332136-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003036129-A1.
20-FEB-2003.
                                                                                                                                               ACF76297 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
                                                                                                                                                                                                                                                                                                                                                                                                                                               Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003104554-A1.
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ACD87836 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
US2003068775-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003049752-A1.
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Best Local Similarity 52.5%; Pred. No. 41;
RESULT 761
                                                                                          Score 38.4; ]
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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cDNA encoding human PRO polypeptide #226.
US2003049765-A1.
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RESULT 764
                                                                                          Match 5.4%;
Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003059879-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                    ACC94289 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027270-A1.
06-FEB-2003.
                                                                                                                                                     Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003040057-A1.
                                                                                                                                                                                                                                                                                                                                                     ACC98024 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003044932-A1.
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     Length 1743;
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                                                          Human secreted/transmembrane protein (PRO) cDNA #226.
US2003040053-A1.
27-FEB-2003.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003032126-A1.
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    Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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US2003054463-A1.
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cDNA encoding human PRO polypeptide #226.
US2003054466-A1.
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    52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
Query Match
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Best Local Similarity
RESULT 778
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RESULT 771
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Best Local Similarity
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Best Local Similarity
RESULT 769
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                                             ACD13104 standard;
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ID NO:451
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068769-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003045700-Al.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068698-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US20003064469-A1.
                                                                 ID NO:451
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ
US2003068760-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003064466-A1.
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                                                ACF01747 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA,
US20031049738-A1.
13-MAR-2003.
 DB 9;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Best Local Similarity 52.5%;
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Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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            Best Local Similarity RESULT 779
                                                                                                                                 Best Local Similarity
RESULT 780
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RESULT 782
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Best Local Similarity
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ACH04508 standard; cDNA; 1743 BP. Human cDNA encoding secreted/transmembrane protein PRO1337.
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10-APR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 801
ID ACD88450 standard; c
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(GETH ) GENENTECH
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RESULT 800
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RESULT 802
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                                                                                                                 Best
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                               ACF28988 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068759-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104552-A1.
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 DB 9; Length 1743;
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                                                                                                                                                                                                                                                                                                                                                                                 DB 9; Length 1743;
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                                                                                                                                                                     ACD90905 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
                                                                                                                                                                                                                                                                                                                 Human secreted/transmembrane protein (PRO) cDNA #226 US2003068765-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003068688-A1.
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                                                                                                                           DB 9;
Score 38.4; I
Pred. No. 41;
                                                                                                                         Score 38.4; I
Pred. No. 41;
                                                                                                                                                                                                                                                   5.4%; Score 38.4; I
52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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cDNA encoding human PRO polypeptide #226.
US2003049754-A1.
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52.5%;
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13-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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           Best Local Similarity
RESULT 788
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RESULT 790
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Best Local Similarity
RESULT 791
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RESULT 794
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Best Local Similarity
RESULT 796
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Best Local Similarity
RESULT 795
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Best Local Similarity
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ACF24018 standard; cDNA; 1743 BP. Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451. US2003068763-A1.
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US2003068739-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003044924-Al.
                                                                                                                                 ACF09939 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
                                                                                      Length 1743;
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ADA78815 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226
US2003073181-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003036126-A1.
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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10-APR-2003
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054465-A1.
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                                                                                 ACC98631 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003044927-A1.
06-MAR-2003.
                                                                                                                                                                                   ACF41936 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003040072-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003040073-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID
US2003064455-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003032124-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003064467-A1.
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Pred. No. 41;
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                                        DB 9;
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Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 815
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
RESULT 809
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Best Local Similarity
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US2003044841-A1.
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ACF23711 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068764-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104551-A1.
                                                                                                                                                                                                                           ACF26225 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068717-A1.
              SEQ ID NO:451
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Human secreted/transmembrane protein (PRO) cDNA #226.
US200304977-A1.
ACF11167 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA,
US2003073170-A1.
                                                                                                       ACF33047 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA,
US2003073176-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003104550-A1.
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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                                                          Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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US2003068728-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 825
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RESULT 821
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Best Local Similarity
RESULT 818
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RESULT 819
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                                                                      Best Local Similarity RESULT 817
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Length 1743;

DB 9;

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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.01520056443-A1.03-APR-2003. (GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US203033178-A1.
                                                                                                                                                                                       Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003064444-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068666-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104538-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US209068774-A1.
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Human secreted/transmembrane protein (PRO) CDNA #226.
US203049781-A1.
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                                                                                                                         Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
           52.5%; Pred. No. 41;
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Pred. No. 41;
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                                          ACF54808 standard; cDNA; 1743 BP
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Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 837
        Best Local Similarity RESULT 835
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RESULT 838
ID ACD49300 standard; c
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Best Local Similarity
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027273-A1.
06-FEB-2003.
                                                                                                                                                                                                                                                                                                                              ACC96294 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
20-8203036161-A1.
20-FEB-2003.
(GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068687-A1.
                                                                                                                       SEQ ID NO:451
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Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451.
US2003059882-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID
US2003059884-A1.
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                                                                                                   ACF10860 standard; cDNA; 1743 BP.

Human secreted polypeptide PRO1337-encoding cDNA, US2003036119-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003036120-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003049739-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003073186-A1.
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 828
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Best Local Similarity
RESULT 832
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RESULT 827
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Best Local Similarity
RESULT 829
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RESULT 845
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RESULT 846
ID ACC9923
DE Human 8
PN US20030
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Query Match

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vuery match
Best Local Similarity 52.5%; Pred. No. 41;
RESULT 863
5.4%; Score 38.4; I
52.5%; Pred. No. 41;
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Best Local Similarity 52.5%;
RESULT 862
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                  (GETH ) GENENTECH INC.
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RESULT 861
            Best Local Similarity RESULT 854
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 Query Match
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Humen secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003073177-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACF79060 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003049764-A1.
                                                                                                                                                                                                                                                                                                                                                                ACF00632 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003054456-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACF14708 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003054457-A1.
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US2003059883-A1.
                                                                                                                                                                                                                                                  ACC99238 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003040067-A1.
                                                                         DB 9; Length 1743;
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Novel human secreted and transmembrane protein PR01337 cDNA.
US2003073129-A1.
                  Human secreted/transmembrane protein (PRO) cDNA #226.
US2003036158-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    #226
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                                                                                                                                                                                                DB 9;
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Human secreted/transmembrane protein (PRO)
US2003054482-A1.
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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                                                                                                                     ACD16941 standard; CDNA; 1743 BP.
CDNA encoding human PRO polypeptide #226.
US2003036151-A1.
     ACD10216 standard; cDNA; 1743 BP.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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BEST LOCAL Similarity PRESULT 853

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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 851
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RESULT 847
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RESULT 848
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20-MAR-2003 Query Match

PN DE

RESULT 849

Query Match

RESULT 852

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Best Local Similarity 52.5%; Score 38.4; DB 9; Length 1743; RESULT 855

ID Acr33661 standard; cDNA; 1743 BP.

BB Human Secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451

PN US2003064450-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068683-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068754-Al.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003104556-A1.
                         SEQ ID NO:451
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Human secreted polypeptide PRO1337-encoding CDNA, SEQ
US200309631-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human secreted/transmembrane protein (PRO) cDNA #226.
US2003068681-A1.
10-APR-2003.
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US2003068731-A1.
ACF51738 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA,
US2003064442-Al.
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Length 1743;

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ACF47133 standard; CDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding CDNA, SEQ ID NO:451.
VS2003068757-A1.
                                                                                                                                              ACF54501 standard; CDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US200308723-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ACF38617 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068766-Al.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068741-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068744-A1.
10-APR-2003.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003104544-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US203068694-Al.
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US2003068715-A1.
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10-APR-2003.
(GETH) GENENTECH INC.
GETY MATCH 5.4%; Score 38.4; D
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
 52.5%; Pred. No. 41;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity RESULT 873
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Best Local Similarity
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                                                                                                   Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003036122-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003054464-A1.
                                                                                                                                                                                                                                                                                                                                                                            ACF06869 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003040065-A1.
27-FEB-2003.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003044928-A1.
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(GETH ) GENENTECH INC.
5.4%; Score 38.4; DB 9; Length 1743;
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          Human secreted/transmembrane protein (PRO) cDNA #226.
US2003049756-A1.
                                                                                                                                                                                                                                 ACUZIO87 standard; CDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
US2001036121-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003040077-A1.
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Pred. No. 41;
                                                                                Score 38.4; DB 9;
Pred. No. 41;
                                                                                                                                                                                              Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
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Human secreted/transmembrane protein (PRO)
US2003064459-A1.
                                                                                                                                                                                                                                                                                                                               5.4%; Score 38.4; I 52.5%; Pred. No. 41;
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Pred. No. 41;
 ACH08595 standard; cDNA; 1743 BP
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                                                                                                                                                                                              5.4%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                              13-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 872
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RESULT 870
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Best Local Similarity
RESULT 871
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Best Local Similarity
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Best Local Similarity
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Length 1743;

DB 9;

Length 1743;

DB 9;

Length 1743;

DB 9;

Length 1743;

DB 9;

ID PN PD

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Query Match
Best Local Similarity
RESULT 892
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                                                                                                                                                                                    Best Local Similarity
RESULT 893
                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 894
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Best Local Si
RESULT 898
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RESULT 882
ID ACF60890 standard; CDNA; 1743 BP.
DE Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
PN US2003036357-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ACF56343 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451.
US2003068708-A1.
                                                                                                                                                                                                                                                                                                                                                               ACF56036 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003068680-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068762-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068761-A1.
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                                                                                      5.4%; Score 38.4; DB 9; Length 1743; 52.5%; Pred. No. 41;
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID
US2003068713-A1.
                                                                                                                                                                                                                                               ADA82886 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
US2003049755-A1.
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Best Local Similarity 52.5%; Pred. No. 41;
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Human secreted/transmembrane protein (PRO) cDNA
US2003054472-A1.
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
                                                                                                                                              cDNA encoding human PRO polypeptide #226.
US2003049760-A1.
                                                                                                                                ACH05777 standard; cDNA; 1743 BP
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(GETH ) GENENTECH INC.
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2003.
2.4 ) GENENTECH IN.
2.TY MATCh
Best Local Similarity 5.
RESULT 887
ID ADB86194 standar
DE Human secret
PN US20030r
PD 20-Y
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                  Best Local Similarity RESULT 883
                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 885
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Best Local Similarity
RESULT 884
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                                                         22-MAY-2003.
(GETH ) GENENTECH
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068771-A1.
                                                                            DB 10; Length 1743;
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Human cDNA encoding secreted/transmembrane protein PRO1337.
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Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003099625-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003054406-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337
US2003083462-A1.
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Human secreted/transmembrane PRO polypeptide cDNA #112.
US2003105013-A1.
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US200308736-Al.
08-MAY-2003
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
5.4%; Score 38.4; D
rry Match
52.5%; Pred. No. 41;
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PA (GETH ) GENENTECH INC.

Query Match

Best Local Similarity 52.5%; Pred. No. 41;

RESULT 895
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Pred. No. 41;
                                                                          5.4%; Score 38.4; I 52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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                                                                                                                                    ADC18104 standard; cDNA; 1743 BP. Human PRO polynucleotide #68. US2003064925-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                             ADD10512 standard; cDNA; 1743 BP.
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RESULT 899
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 897
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Length 1743;

Length 1743;

Length 1743;

Length 1743;

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DB 10; Length 1743;
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                                                                                   ADG02919 standard; cDNA; 1743 BP.
Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003207397-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2002207392-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA US2003207399-A1.
06-NOV-2003.
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Human cDNA encoding secreted/transmembrane protein PRO1337
US2003065142-A1.
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                                       Score 38.4; ]
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Human PRO polynucleotide #226.
US2003027272-Al.
                                       52.5%;
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Best Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                      Query Match
Best Local Similarity
                                                                                                                                                                                    Best Local Similarity RESULT 911
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Best Local Similarity
RESULT 916
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ID ACA71205 standard;
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                                                    DB 10; Length 1743;
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                                                                                             ADD39350 standard; cDNA; 1743 BP.
Human cDNA encoding secreted/transmembrane protein PRO1337
US2003096954-A1.
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RESULT 905
ID ADE20137 standard; cDNA, 1743 BP.
DE Human cDNA encoding secreted/transmembrane protein PRO1337.
PN US2003092883-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337,
US2003204054-A1.
                                                                                                                                                                                                                                                 Human cDNA encoding secreted/transmembrane protein PRO1337 V820039925061-Al. 15-MAY-2003 (GETH ) GENENTECH INC.
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Human cDNA encoding secreted/transmembrane protein PRO1337
US2003082628-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003204053-A1,
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                                                                                                                                                                                    DB 10;
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                                                  Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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             22-MAY-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 903
                                                                Best Local Similarity
RESULT 901
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Best Local Similarity
RESULT 904
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Best Local Similarity
RESULT 908
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US2003096955-A1.
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27-FEB-2003.
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                 Query Match
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003040056-A1.
ACC87733 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027278-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ACC89575 standard; cDNA; 1743 BP. Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003027264-A1. 6-FEB-2003.
                                                                                                                      ACC87119 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003036159-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003036144-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003036147-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
27-FEB-2003
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003044923-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
20.010303149-A1.
20-FEB-2003.
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Human secreted/transmembrane protein (PRO) cDNA #226.
222033040070-A1.
27-FBB-2003.
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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cDNA encoding human PRO polypeptide #226.
US2003032113-A1.
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Best Local Similarity
RESULT 920
ID ACC87119 stand
DE Human Ser
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Best Local Similarity
RESULT 929
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Best Local Similarity
RESULT 927
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Best Local Similarity
RESULT 922
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Best Local Similarity
RESULT 923
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Best Local Similarity
RESULT 926
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Best Local Similarity
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SEQ ID NO:451
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ACA92175 standard; cDNA; 1743 BP.
Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003027277-A1.
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Novel human secreted and transmembrane protein PR01337 cDNA.
US2003073813-A1.
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Novel human secreted and transmembrane protein PR01337 cDNA.
US2003013153-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003211572-A1.
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Pred. No. 41;
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Pred. No. 41;
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 DB 10;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
 Score 38.4; I
Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Human PRO polynucleotide #226.
US2003036140-A1.
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Best Local Similarity 52.5%;
RESULT 938
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Best Local Similarity 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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            Best Local Similarity RESULT 930
                                                                                                                    Best Local Similarity RESULT 931
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5.4%; Score 38.4; DB 12; Length 1743; 52.5%; Pred. No. 41;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003068770-A1.
                                                                                                                                                                                                                                                                                                                             ADH03600 standard; cDNA; 1743 BP.

Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003224478-A1.
04-DEC-2003.
CERT ) GENENTECH INC.
                                         AUH03123 standard; CDNA; 1743 BP.
Human CDNA encoding secreted/transmembrane protein PRO1337.
002003216562-A1.
                                                                                                                                                                                           ADH04077 standard, cDNA, 1743 BP.
Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003220471-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADH04554 standard; CDNA; 1743 BP.
Human CDNA encoding secreted/transmembrane protein PRO1337.
US2004005626-A1.
08-JAN-2004.
(GETH ) GENENTECH INC.
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                                                                                                                                              Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
5.4%; Score 38.4;
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Human PRO polynucleotide #226.
US2003068768-A1.
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Human PRO polynucleotide #226.
US2004023321-A1.
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Best Local Similarity 52.5%;
                                                                                                       20-NOV-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                                            Query Match
Best Local Similarity
RESULT 950
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 Query Match
Best Local Similarity
RESULT 949
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Best Local Similarity
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Best Local Similarity
RESULT 957
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                                    DB 12; Length 1743;
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Novel human secreted and transmembrane protein PRO1337 cDNA, 10S200321590-A1.
20-NOV-2003.
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Novel human secreted and transmembrane protein PRO1337 cDNA 10S209215912-A1.
20-NOV-2003.
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Novel human secreted and transmembrane protein PRO1337 cDNA
US2003215911-A1.
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Human CDNA encoding secreted/transmembrane protein PRO1337.
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Human cDNA encoding secreted/transmembrane protein PRO1337.
US2003195334-A1.
                                                                                  ADE96608 standard; cDNA; 1743 BP.
Human cDNA encoding secreted/transmembrane protein PRO1337
US2003195347-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337
US2003199675-A1.
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                                  5.4%; Score 38.4; I 52.5%; Pred. No. 41;
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                                                                                                                                                                         52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 942
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                                               Best Local Similarity
RESULT 940
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Best Local Similarity
RESULT 941
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Best Local Similarity
RESULT 943
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Best Local Similarity
RESULT 944
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US2003211574-A1
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Best RESULT

Query Match

RESULT

Query Match

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Query Match

RESULT 964

Query Match

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ADG79359 standard; cDNA; 3337 BP.
Human secreted protein cDNA of the invention SEQ ID NO:165.
WO200268638-A1.
06-SEP-2002.
(HUMA-) HUMAN GENOME SCI INC.
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US2002160410-A1.
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58.3%;
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PA (EPIG-) EPIGENOMICS AG.
Query Match 5.4%;
Best Local Similarity 45.2%;
RESULT 972
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
Query Match 5
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(HADL/) HADLACZKY G.
(SZAL/) SZALAY A A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Ouery Match
Best Local Similarity
RESULT 976
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Best Local Similarity
RESULT 969
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                                                                                                                                     5.4%; Score 38.4; DB 12; Length 1743; 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match 5.4%; Score 38.4; DB 12; Length 1743;
Best Local Similarity 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                        Length 1743;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2004048334-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2004053358-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2004049335-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1337.
US200407315-A1.
15-APR-2004.
(GETH ) GENENTECH INC.
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Arabidopsis thaliana DNA fragment SEQ ID NO: 49004.
EP1033405-A2.
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52.5%; Pred. No. 41;
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         52.5%; Pred. No. 41;
                                                                                                                                                                                               ADJ64860 standard; cDNA; 1743 BP.
Human PRO polynucleotide #226.
US2004038337-A1.
                                               ADK83001 standard; cDNA; 1743 BP
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                                                              Human PRO polymucleotide #112
VSZO04404927-A1.
04-MAR-2004.
(GETH ) GENENTECH INC.
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PA (GETH ) GENENTECH INC.
QUETY MATCh
Best Local Similarity 52.5%;
RESULT 960
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(BEHW ) BEHRINGWERKE AG.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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P.falciparum GBP130h.
EP499834-A2.
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Best Local Similarity
RESULT 963
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Best Local Similarity
RESULT 967
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       Best Local Similarity
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DB 6; Length 37515;
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                                                                                                                                                                 Length 4010;
                                                                                                                                                                                                                                                                                                                            DB 6; Length 6185;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       6; Length 6880;
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          DB 6; Length 3337;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAS61223 standard; DNA; 6880 BP.
Human gene regulation-associated gene oligonucleotide #178
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABQ66997 standard; DNA; 37515 BP.
Human angiogenesis associated polynucleotide SEQ ID NO 27.
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human chemically pretreated gene sequence #77 strand WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                          #83
                                                                  AAL26509 standard; cDNA, 4010 BP.
Human breast cancer expressed polynucleotide 18966.
19-JUL-2001.
MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.4%; Score 38.4; DB 4; Le
Ft Local Similarity 50.0%; Pred. No. 49;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADD/0293 standard; DNA; 6880 BP. Chemically treated cell signalling DNA sequence#92. W0200202807-A2.
                                                                                                                                                                                                                  ABL32413 standard; DNA; 6185 BP.
Human immune system associated gene SEQ ID NO: 386.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                        Signal transduction associated gene modified DNA : w02002028-A2. 03-UAN-2002. (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 6;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         wuery match 5.4%; Score 38.4; I Best Local Similarity 47.5%; Pred. No. 59; RESULT 974
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 38.4; 1
Pred. No. 78;
Vuely Match
Beet Local Similarity 50.0%; Pred. No. 47;
RESULT 968
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     18-OCT-2001.
(EPPL) EPIGENOMICS AG.
5-4%; Score 38.4; I
St Local Similarity 45.2%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 38.4; I
Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 38.4;
Pred. No. 70;
                                                                                                                                                                                                                                                                                                                            5.4%; Score 38.4; 1 55.1%; Pred. No. 54;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 38.4; Pred. No. 55;
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ID DE PN PD PD PA PA PA PA PA PA

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ID ADATIOS standard; DNA; 2000 BP.

DE Rice gene, SEQ ID 5263.

PN W02003000898-Al.

PD 03-JAN-2003.

PA (SYGN ) SYNGENTA PARTICIPATIONS AG.

Query Match

Some 38.2;

Best Local Similarity 8.4%; Pred. No. 47;

RESULT 986
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.4%;
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(EXSE-) EXSEED GENETICS LLC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABL33760 standard; DNA; 7851
                                                                                                                                                                                                                                                                                                                                                                               TWINE N C.
BURCZYNSKI M E.
TREPICCHIO W L.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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Best Local Similarity
                                                                                                                                                                                                                                                                                               ADP13500 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity
RESULT 991
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10-APR-2003.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                               (AMHP ) WYETH.
                                                                                                                                                                                                                                                                                                                                                 10-JUN-2004
                                                                                                                                                                                                                                                                                                                                                                               (TWIN/)
(BURC/)
(TREP/)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                (DORN/)
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          RESULT 985
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 989
                                                                                                                                                                                                                                                               Best |
RESULT
                                                                                                                                                                                                                                                                                                                ABQ41663 standard; DNA; 1164 BP.
Oligonuclectide for detecting cytosine methylation SBQ ID NO 28254.
WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABQ41662 standard; DNA; 1164 BP. Oligonucleotide for detecting cytosine methylation SEQ ID NO 28253. WO200218632-A2.
AAV21210 standard; DNA; 58407 BP.
Methanococcus jannaschii large circular extrachromosomal element.
W09807830-A2.
26-FEB-1998.
(GENO-) INST GENOMIC RES.
(UNII ) UNIV ILLINOIS FOUND.
(UNII ) UNIV JOHNS HOPKINS SCHOOL MEDICINE.
3TY MATCh
3TY MATCh
3TY COOL SIMILARILY 47.5%; Pred. NO. 86;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAV70954 standard; DNA; 1141 BP.
Zea mays 10 kDA zein gene portion used as a promoter in plasmids
WO9844780-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                       ABS05479 standard; DNA; 456 BP.
Human genome-derived single exon probe from lung SEQ ID No 5470.
WO200186003-A2.
                                                                                                                                                                                                                                                        Length 177851;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
(ery Match 5.4%; Score 38.2; DB 12; Length 1486;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 1141;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 456
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH34164 standard; cDNA; 511 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:1246
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human soft tissue sarcoma-upregulated DNA - SEQ ID 5536 WO2004048938-A2.
                                                                                                                                                                                                                                                        5.4%; Score 38.4; DB 8; 56.2%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                   27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     15-NOV-2001.

(MOLE-) MOLECULAR DYNAMICS INC.

(MOLE-) MAtch

5.4%; Score 38.2; DB 6;

ery Match

cry Match

5.4%; Pred. No. 34;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Wolver-201.
05-AP-2001.
(HUMA-) HUMAN GENOME SCI INC.
Lery Match 58.3%; Score 38.2; DB 4;
Lery Match 14 58.3%; Pred. No. 35;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.4%; Score 38.2; DB 2; 53.7%; Pred. No. 42;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              07-MAC-2002.
(EPIG-) EPIGENOMICS AG.
(ETY MAtch 5.4%; Score 38.2; DB 6;
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                                                                                                                                                                                         DNA
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                                                                                                                                                                 AAL57272 standard, DNA; 177851 BP.
bA438B23-1 'human modifier of p53 pathway'
                                                                                                                                                                                                                                                                                                                                                                                           52.1%; Score 38.2; I 52.1%; Pred. No. 32;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 38.2;
Pred. No. 42;
                                                                                                                                                                                                                                                                                                      ADL43989 standard; DNA; 324 BP.
Human ovarian cancer DNA marker #17879.
WO200170979-A2.
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07-MAR-2002.
(EPIG-) EPIGENOMICS AG.
... Match :1arity 52.9%;
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(EXSE-) EXSEED GENETICS LLC.
                                                                                                                                                                                                                                        (EXEL-) EXELIXIS INC.
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Best Local Similarity
RESULT 979
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Best Local Similarity
RESULT 982
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Best Local Similarity
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WO2003035833-A2.
01-MAY-2003.
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Best Local Si
RESULT 977
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DB 12; Length 2623;
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DB 8; Length 2000;
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Human chemically modified disease associated gene SEQ ID NO
WO200200927-A2.
                                                                                                                                                                                                                 Renal cell carcinoma differentially expressed gene #236.
WO2004048933-A2.
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WO200200928-A2.
03-JAN-2002.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human prostate expression marker cDNA 30190.
W0200160860-A2.
                                                                                                                                              Score 38.2; Pred. No. 50;
                                                                                                                                                                                                                                                                                                                                                                                                     5.4%; Score 38.2; I
45.4%; Pred. No. 50;
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Pred. No. 55;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC
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Pred. No. 56;
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Best Local Similarity 47.8%; Pred. No. 63;
RESULT 992
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Pred. No. 72;
                                                                                                                                                                                                                                                                                                                                                                                                                                         Human novel cDNA; 4414 BP.
Human novel cDNA sequence, SEQ ID NO:776.
10-abb-201
                                                                    Zea mays 10 kDA zein gene DNA sequence. WO9844780-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          25-APR-2002.
(CALY ) CALIFORNIA INST OF TECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABK52611 standard; DNA; 4755 BP.
DNA encoding human Claspin protein.
WO200233115-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL32672 standard; DNA; 16287 BP.
                                                 AAV70953 standard; DNA; 2562 BP.
                                                                                                                                                                                                 DNA; 2623 BP
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06-NOV-2003
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Primary rat hepatocyte toxicity modelling related gene SEQ ID NO:2454.
WQ2003065993-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADG33178 standard; DNA; 1381 BP.
Human DNA differentially expressed in patients with SLE SegID502.
WO2003090694-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 200620;
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                                                                                                                                                                                                                                                                                                                         Length 119950;
                                                    DB 6; Length 16287;
                                                                                                                                                                                        DB 6; Length 58837;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human presynaptic cytomatrix protein, REP82, genomic sequence WO200404164-A2.
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Best Local Similarity 62.8%; Pred. No. 39;
RESULT 1000
ID AAS62239 standard; CDNA; 681 BP.
DE CDNA sequence #26 encoding novel human secreted protein.
PN WO200177291-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            "wery March Best Local Similarity 54.7%; Pred. No. 1.26+02; RESULT 998
ID AAI82415 standard; CDNA; 447 BP. DE Human polymucleotide SEQ In "P. Prog. No. 07-SEP-20"
PM WOO10164835-A2.
PA (""")
                                                                                                                                                                                                                                                                                                                                                                          ADL13909 standard; DNA; 129588 BP. Osteoarthritis-associated polymorphic nucleotide #441. WO2003054166-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                          PA (INCY-) INCYTE GENOMICS INC.
Querry Match
5.4%; Score 38.2; DB 10;
Best Local Similarity 63.7%; Pred. No. 1.1e+02;
RESULT 997
                                                                                                                                                                                                                                                                                                                            5.4%; Score 38.2; DB 2; 51.5%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PD 14-AUG-2003.
PA (GBNE-) GENE LOGIC INC.
Query Match
Best Local Similarity 57.6%; Pred. No. 44;
RESULT 1002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB 6;
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                                                      5.4%; Score 38.2; I
47.0%; Pred. No. 73;
                                                                                                                                                                                        58.3%; Pred. No. 96;
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(REHO-) CENT RECH HOPITAL SAINTE-JUSTINE.
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                                                                                                                                                         ) 25-APR-2002.

(CALY ) CALIFORNIA INST OF TECHNOLOGY.

Query Match 5.4%; Score 38.
                                                                                                                                                                                                                                           AAX90201 standard; DNA; 119950 BP
                                                                                                  ABK52612 standard, DNA; 58837 BP. Human Claspin genomic sequence. WO200233115-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAT42808 standard; DNA; 486 BP.
Polymorphic locus Q900 sequence.
WO9634979-A2.
                                                                                                                                                                                                                                                                                                           (MILL-) MILLENNIUM PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          18-OCT-2001.
(GEMY ) GENETICS INST INC.
                     03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                  LOO2.

LOO2.

LOO2.

LOO3.

LY Match

Best Local Similarity

RESULT 995

ID AAX90201 stand-
DE Human yes1

PN WC99357
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                                                             Best Local Similarity
RESULT 994
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Best Local Similarity
RESULT 999
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       WO200200928-A2.
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                                                      Query Match
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ABL70146 standard; DNA; 6754 BP.
Chemically treated cell signalling DNA sequence complementary to#118.
WO200202807-A2.
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Human tyrosine phosphatase SHP1 bisulphited genomic DNA SeqID JP2004000128-A.
08-JAN-20040
             Length 1381;
                                                                                                                                          Length 1755;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PD 10-JAN-2002.

PA (EPIG-) EPIGENOMICS AG.

S.3%; Score 38; DB 6; Length 6754;

Best Local Similarity 47.7%; Pred. No. 67;

RESULT 1008

ID AAS61305 standard; DNA; 6754 BP.

DE Human gene regulation-associated gene oligonucleotide #260.

PN W0200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 6754;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 9810;
                                                                                                                                                                                                                                                                                                                                                                                                        Length 5461;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 7615
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 5461
                                                                                                                                                                                                                                                                         Length 3067
                                                                                                                                                                                          ADG79262 standard; cDNA; 3067 BP.
Human secreted protein cDNA of the invention SEQ ID NO:68.
WO200268638-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     7.
                                                                                                                                                                                                                                                                                                                                                                                                                                      ABQ66977 standard; DNA; 5461 BP.
Human angiogenesis associated polynucleotide SEQ ID NO
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human chemically pretreated gene sequence #27 strand WO200202806-A2.
                                                                             #978
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL32426 standard; DNA; 9810 BP.
Human immune system associated gene SBQ ID NO: 399.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                         971
                                                                             Π
               DB 10;
                                                                                                                                            DB 12;
51;
                                                                                                                                                                                                                                                                                                                                    Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 6;
69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 6;
73;
                                                                                                                                                                                                                                                                         DB 6;
57;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 6;
67;
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9
                                                            ADP28980 standard; DNA; 1755 BP.
Human secreted protein encoding sequence SEQ
Human secreted protein encoding sequence SEQ
29-APR-2004.
(FIVE) FIVE PRIME THERAPEUTICS INC.
(FIVE) 5.3%; Score 38; DB 12
ety Match
the Local Similarity 62.8%; Pred. No. 51;
                                                                                                                                                                                                                                                                                                                                                                                                        DB
65;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB
65;
                                                                                                                                                                                                                                                                                                                                                                                                        Score 38;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 38;
Pred. No.
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Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 38;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 38;
Pred. No.
             5.3%; Score 38; 51.8%; Pred. No.
                                                                                                                                                                                                                                              06-SEP-2002.
(HUMA-) HUMAN GENOME SCI INC.
EXY MATCH 5.3%; SCORE 38;
St Local Similarity 49.2%; Fred. No.
A (EXPR-) EXPRESSION DIAGNOSTICS INC.
Query Match 5.3%; Score
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABK39972 standard; DNA; 7615 BP.
                                                                                                                                                                                                                                                                                            Best Local Simitarity Town RESULT 1005
ID ABL32998 standard, DNA, 5461 BP.
                                                                                                                                                                                                                                                                                                                                                              Word-2002.
(BPIG-) EPIGENOMICS AG.
FPIG-) Adich (STik)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
sry Match 5.3%;
st Local Similarity 53.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5.3%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
Query Match 5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
                            Best Local Similarity RESULT 1003
                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1007
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 1009
                                                                                                                                                           Best Local Similarity RESULT 1004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                              Query Match
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                                                                                                                                               Query Match
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us-09-989-293a-376.rng.spdi

03-JAN-200

Query Match

Query Match

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Probe #701 for gene expression analysis in human heart cell sample. WO200157274-A2.
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ID ABQ46793 standard; DNA; 538 BP.
DE Oligonucleotide for detecting cytosine methylation SEQ ID NO 33384.
PN W0200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABQ46792 standard, DNA, 538 BP. Oligonucleotide for detecting cytosine methylation SEQ ID NO 33383. WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAI00717 standard; DNA; 470 BP.
Probe #708 used to measure gene expression in human breast sample.
WO200157270-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABSO0747 standard; DNA; 470 BP.
Human genome-derived single exon probe from lung SEQ ID No 738
WO200186003-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.3%; Score 37.8; DB 4; Length 1160;
                                                                                                                                                                                                                                                      719
          5.3%; Score 37.8; DB 4; Length 470; 53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                  Length 470;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 470;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 6; Length 538;
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                                                                                                                                                                                                                                                    NO.
                                                                                                                                                                         Length
                                                                                                                                                                                                                           AAK26162 standard; DNA; 470 BP.
Human bone marrow expressed single exon probe SEQ ID
WO200157276-A2.
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ID AAL26474 standard; cDNA; 1160 BP.

DE Human breast cancer expressed polynucleotide 18931.

PN W0200151628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           4,
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                                                                                                                                                                     DB 4;
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                                                                                                                                                                                                                                                                                                                                                                                    AAK00710 standard; DNA; 470 BP.
Human brain expressed single exon probe SEQ ID
WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABS25751 standard; DNA; 470 BP.
Human liver single exon probe, SEQ ID No 741.
WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB
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                                                                                                                 MOJECTORY OF AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
5.3%; Score 37.8; D
tery Match
5.3%; Pred. No. 43;
                                                                                                                                                                                                                                                                            MOST OF AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
5.3%; Score 37.8; I
cery Match 53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                              09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
5.3%; Seore 37.8; Ist Local Similarity 53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5.3%; Score 37.8; 1
53.8%; Pred. No. 43;
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53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           IS-NOV-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
Ery Match
Strical Similarity 53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 37.8; I
Pred. No. 44;
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Pred. No. 44;
                                                                      DNA; 470 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         WCACALON ON WAS CALLED OF BEIGHOMICS AG. (BPIG-) EPIGENOMICS AG. 5.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  52.9%;
                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
  Query Match
Best Local Similarity
RESULT 1021
                                                              ABA22235 standard;
                                                                                                                                                                   Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SEQ ID NO:15.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAI10792 standard; DNA; 470 BP.
Probe #725 for gene expression analysis in human cervical cell sample.
WO200157278-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                           45.
                                                                                                                                                                                                                                                                                                                                                                                      ABL49345 standard; DNA; 19233 BP.
Human polynucleotide associated with DNA replication SEQ ID NO
W0200177377-A2.
                                                                                                                                                                                                                                   ABL30294 standard; DNA; 18396 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 42355.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 266145
                    5.3%; Score 38; DB 12; Length 10427; 51.1%; Pred. No. 74;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AA132050 standard; DNA; 470 BP.
Probe #736 used to measure gene expression in human placenta
WO200157272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (KYOW ) KYOWA HAKKO KOGYO KK.

(KYOW ) KYOWA HAKKO KOGYO KK.

5.3%; Score 38; DB 4; Length 80578;

ery Match

55.2%; Pred. No. 1.1e+02;
                                                                                                                                                                                   Length 15548,
                                                                                                                                                                                                                                                                                                                                             Length 18396;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 19233,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.3%; Score 37.8; DB 4; Length 470, 53.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 470
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAH44800 standard; DNA; 80578 BP.
Human GPCR protein KAT06734L DNA containing exons 1 and 2
JP2001245666-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    #747
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ABA42021 standard, DNA, 470 BP.
Human breast cell single exon nucleic acid probe #716.
WO200157271-A2.
                                                                            ABL34155 standard; DNA; 15548 BP.
Human immune system associated gene SEQ ID NO: 2128
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABA52442 standard; DNA; 470 BP.
Human foetal liver single exon nucleic acid probe
WO200157277-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(ery Match 5.3%; Score 37.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  WOLUSS...
12-JUN-2003.
(1215) ISIS INNOVATION LTD.
(1217 Match 5.3%; Score 38; DB 10;
Ery Match 6.0%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                          DB 4;
83;
                                                                                                                                            03-JAN-ZUUG.
(EPIG-) EPIGENOMICS AG.
:ry Match 5.3%; Score 38; DB 6;
-- Tanal Similarity 47.2%; Pred. No. 80;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.3%; Score 38; DB 6; 53.6%; Pred. No. 84;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (MOLE-) MOLECULAR DYNAMICS INC.
ry Match
t Local Similarity 53.8%; Pred. No. 43;
(KAGA-) KAGAKU GIJUTSU SHINKO JIGYODAN.
                                                                                                                                                                                                                                                                                                                                        5.3%; Score 38; 51.8%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADE87477 standard; DNA; 266145 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (MOLE-) MOLECULAR DYNAMICS INC.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Fowlpox virus genome DNA. WO2003047617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                      18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 1018
                                   Best Local Similarity
RESULT 1012
                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1013
                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                RESULT 1014
ID ABL49345 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1015
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1016
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Best Local Similarity
RESULT 1017
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RESULT 1019
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09-AUG-2003

Query Match

Query Match

09-AUG-200

09-AUG-2003

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03-JAN-2002
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ABK31471 stansduction associated gene modified complementary DNA #157.
WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.3%; Score 37.8; DB 12; Length 3953; 49.8%; Pred. No. 67;
                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 37.8; DB 12; Length 2000;
Pred. No. 58;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 3953;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 6242;
                                                                                                                                                                                                 Length 1811;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 6731;
                                                                                              DB 4; Length 1537;
                                                                                                                           AAS21090 standard; cDNA; 1811 BP.
cDNA encoding human ubiquitin-conjugating enzyme 10_01
WO200194407-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABK39963 standard; DNA; 6731 BP. Human chemically pretreated gene sequence #22 strand WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABL34148 standard; DNA; 6242 BP.
Human immune system associated gene SBQ ID NO: 2121.
W0200200928-A2.
(BPIGS)
BPIGENOMICS AG.
5.38; Score 37.8; DB 6; Lerery Match
Social Similarity 50.2%; Pred. No. 74;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.3%; Score 37.8; DB 6; 49.8%; Pred. No. 67;
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9
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10-JANTZOUZ.

5.3%; SCOTE 37.8; DB

PROPERTY MATCH
1.7-APA | Similarity 48.4%; Pred. No. 75;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB
                                                                                                                                                                         (SHAN-) SHANGHAI BIOWINDOW GENE DEV INC.
SHAN-) SHANGHAI BIOWINDOW GENE DEV INC.
5.3%; Score 37.8; Inc. 1.00.8 Similarity 64.0%; Pred. No. 57;
                                                                                              5.3%; Score 37.8; I
45.0%; Pred. No. 55;
     49,8%; Pred. No. 52;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABZ11664 standard; cDNA; 3953 BP.
Human polynucleotide SEQ ID NO 546
WO200270539-A2.
                                                                                                                                                                                                                                         ADJ41453 standard; cDNA; 2000 BP
                           AAF58662 standard; DNA; 1537 BP.
Porcine myostatin gene promoter.
BP1072680-A1.
31-JAN-2001.
(PPIZ ) PPIZER PROD INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                              5.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  BUDWORTH P.
MOUGHAMER T.
BRIGGS S P.
COOPER B.
GLAZEBROOK J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 1037
ID ABK31471 standard; DN
DE Signal transduction PN WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (XUEA/) XUE A.
(DRMA/) DRMANAC R T.
                                                                                                Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1035
                                                                                                                                                                                                                                                                                                                                                                GOFF S A.
KATAGIRI F.
                                                                                                                                                                                                                                                                                                                                                                                          (KREP/) KREPS J.
(PROV/) PROVART N.
(RICK/) RICKE D.
(ZHUT/) ZHU T.
    Best Local Similarity
                                                                                                                                                                                                     Query Match
Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                          Plant cDNA #2453.
US2004016025-A1.
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                                                                                                                                                                                                                                                                                                 (BUDM/)
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RESULT 1033
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                   RESULT
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ABZ09992 standard; DNA; 7369 BP.
Haematopoietic cell proliferation disorder related DNA sequence #132.
WO200277272-A2.
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Haemacopoietic cell proliferation disorder related DNA sequence #278.
WO200277272-A2.
03-OCT-2002.
                                                                                     to#228.
                                                                        DNA; 7072 BP. cell signalling DNA sequence complementary
                                                                                                                                                                                                                                                                                                                                                                                                                         DB 8; Length 7369;
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PD 20-FEB-2003.

Query Match 5.3%; Score 37.8; DB 8; Length 7369;

Best Local Similarity 51.5%; Pred. No. 76;

RESULT 1043

DE Human 5' and/or regulatory region of CDKNIA DNA SEQ ID NO:19.

PN WO200277272-A2.
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                      Length 7072;
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                                                                                                                                                   Length 7072
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Human gene regulation-associated gene oligonucleotide #340.
W0200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ACF62808 standard, DNA, 7369 BP.
Colon cancer analysis related genomic DNA SEQ ID NO:57.
WO2003014388-A2.
                                                                                                                                                                                                                                                                                                                                                      Colon cancer analysis related genomic DNA SEQ ID NO:35 WO2003014388-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 8;
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                        DB 6;
                                                                                                                                                       DB 6;
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DNA transcription associated genomic DNA #141.
WO200192565-A2.
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                                                                                                                                                                                                                                                                             vuery Match 5.3%; Score 37.8; I
Best Local Similarity 50.3%; Pred. No. 76;
RESULT 1040
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Pred. No. 76;
U3-UMN-2002.
(EPIG-) EPIGENOMICS AG.
127 Match
17 Incal Similarity 50.3%; Pred. No. 76;
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Pred. No. 76;
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Pred. No. 76;
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Pred. No. 76;
                                                                                                                                                     Score 37.8; 1
Pred. No. 76;
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Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human p21 genomic DNA; 7369 BP.

### PACO GENOMIC DNA SEQ ID NO:2.

### PACO GENOMIC DNA SEQ ID NO:2.
                                                                                                                                                                                                                                                                                                                                         ACF62786 standard; DNA; 7369 BP.
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(EPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
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03-OCT-2002.
(EPIG-) EPIGENOMICS AG.
5.3$;
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RESULT 1045
                                                                                                                  Mvcv.
(BPIG-) EPIGENOMICS AG.
ery Match 5.3%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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                      Query Match
Best Local Similarity
RESULT 1038
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Best Local Similarity
RESULT 1042
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RESULT 1044
                                                                                                                                                                      Best Local Similarity RESULT 1039
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                                                                       ABL70566 standard;
Chemically treated
WO200202807-A2.
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ABL70604 standard; DNA; 34548 BP. Chemically treated cell signalling DNA sequence complementary to#247. WO200202807-A2.
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5.3%; Score 37.8; DB 10; Length 202001;
51.5%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                 Match 5.3%; Score 37.8; DB 10; Length 29956; Local Similarity 58.4%; Pred. No. 1e+02;
                                                                                                                                                                       10; Length 29956;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match 5.3%; Score 37.8; DB 12; Length 29956; Best Local Similarity 58.4%; Pred. No. 1e+02;
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      5.3%; Score 37.8; DB 9; Length 29956; 58.4%; Pred. No. 1e+02;
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ID AAN90224 standard; DNA; 1052 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 306;
                                                                                                                                                                                                                                                                                                                                                                            AUM/4558 standard; DNA; 29956 BP.
Murine carcinoma associated (CA) nucleic acid #115.
15.2004072154-Al.
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 5.3%; Score 37.6; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.3%; Score 37.8; DB 6; 53.8%; Pred. No. 1.3e+02;
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Tansporter protein genomic DNA.

Query Match

5.3%; Score 37.8; DB 6;

Best Local Similarity 51.5%; Pred, No. 1.5e+02;
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Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 37.8; DB 6; Pred. No. 1.3e+02;
                                                                                                                      PD 30-2030.

PA (SAGR-) SAGRES DISCOVERY.

Query Match
Best Local Similarity 58.4%; Pred. No. 1e+02,

RESULT 1057

ID ADCRS443 standard; DNA, 29956 BP.

DE Mouse Lick genomic sequence.

PN W02003045230-A2.
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Leukaemia-related DNA sequence #2812.
WO2003039443-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (DEKR-) DEUT KREBSFORSCHUNGSZENTRUM. (UYLU-) UNIV LUDWIG MAXIMILIANS. (HAFE/) HAFERLACH T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADG46742 standard; DNA; 202001 BP. Human transporter genomic DNA.
                                                              ADB72701 standard; DNA; 29956 BP
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.3%;
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(SAGR-) SAGRES DISCOVERY
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(EPIG-) EPIGENOMICS AG.
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(ENGE/) ENGELHARD E K.
      Query Match
Best Local Similarity
RESULT 1056
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
                                                                                   Mouse Lck gene.
W02003008583-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                              15-APR-2004
                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                        ABK31249 standard; DNA; 10543 BP. Signal transduction associated gene modified complementary DNA #46. WO200200926-A2.
                                                                             AAS46304 standard; DNA; 10369 BP.
Tumour suppressor gene derived chemically modified sequence #26
WO200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABN80146 standard; DNA; 14920 BP.
Human chemically modified disease associated gene SEQ ID NO 163
WO200200927-A2.
                                                                                                                                                                                    DB 4; Length 10369;
                                                                                                                                                                                                                                                                                                                                                       Length 10369;
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                     DB 6; Length 9504;
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Human immune system associated gene SEQ ID NO: 1590
WO200200928-A2.
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Mouse Lck carcinoma associated gene, SEQ ID NO:1481.
W02003057146-A2.
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WO200200928-A2.
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03-JAN-2002.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.3%; Score 37.8; DB 6;
tery Match
51.5%; Pred. No. 82;
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WOLD STORM STORY (EPTG.) BPIGENOMICS AG.
(EPTG.) EPIGENOMICS AG.
(EPT Match 5.3%; Score 37.8; DB 6; CTY Match 7.3%; Pred. No. 82;
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Human immune system associated gene SEQ ID NO:
WO200200928-A2.
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18-OCT-2001.
18-OCT-2001.
18-OCT-2001.
19-OCT-2001.
5.3%; Score 37.8; DB tery Match 5.3%; Pred. No. 82;
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20-ESP-2001.
20-EPIG-) BPIGENOMICS AG.
5.3%; Score 37.8; D
1ery Match
51.5%; Pred. No. 82;
                 5.3%; Score 37.8; I
57.0%; Pred. No. 81;
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44.0%; Pred. No. 91;
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48.8%; Pred. No. 94;
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                                      57.0%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
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Best Local Similarity
RESULT 1048
                                  Best Local Similarity
RESULT 1047
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RESULT 1050
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RESULT 1052
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RESULT 1053
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Best Local Similarity
RESULT 1055
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RESULT 1054
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Query Match
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                                                                                                                                                                                                                                                                                                                                     AAA70123 standard; DNA; 1452 BP.
Plasmodium falciparum chromosome 2 related DNA sequence SEQ ID NO:256.
MC20025728-A2.
11.MAY-2000.
(HOFF) HOFFMAN S.
(CARU), CARUCCI D.
(GARD), SARDNER M.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQ ID NO:22
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 1984;
                                                                                                                                                                              DB 12; Length 1140;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 37.6; DB 6; Length 6240; Pred. No. 82;
                                                                                                                                                                                                                                                                                                Length 1335;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 6149;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 5218;
                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 3; Length 1452;
                                                          Length 1052;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADE37776 standard; DNA; 1984 BP.
Human chemically treated calcitonin nucleotide seguence
WO2003072820-A2.
                                                                                                        ADQ23230 standard; DNA; 1140 BP.
Human soft Lissue sarcoma-upregulated DNA - SEQ ID 6050
W02004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABL33266 standard, DNA, 5218 BP.
Human immune system associated gene SEQ ID NO: 1239
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            #1
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Pred. No. 65;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAS63306 standard; DNA; 6149 BP.
Chemically pretreated metabolism associated gene
W02017451.A2.
18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                 DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     . 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.3%; Score 37.6; DB 6; 49.5%; Pred. No. 79;
                                                             DB 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABLJ32049 standard; DNA; 6240 BP.
Human immune system associated gene SEQ ID NO:
WQZ0Q220928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 37.6; DB
Pred. No. 82;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        04-SEF-zvus.
(EPIG-) EPIGENOMICS AG.
sry Match
** Toral Similarity 51.2%; Pred. No. 65;
                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.3%; Score 37.6; I
56.2%; Pred. No. 60;
                                                                                                                                                                (PROT-) PROTEIN DESIGN LABS INC.
ry Match
5.3%; Score 37.6; I
t Local Similarity 55.3%; Pred. No. 58;
                                                          Score 37.6; 1
Pred. No. 57;
                                                                                                                                                                                                                                                                                (GENO-) GENOME THERAPEUTICS CORP.

ry Match
5.3%; Score 37.6;
t Local Similarity 49.5%; Pred. No. 59;
Malaria-specific DNA insert of clone 41-2.
EP322712-A.
05-JUL-1989.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Pretreated genomic DNA region 227 W02003072821-A2.
                                                                                                                                                                                                                        Aproli65 standard; DNA; 1335 BP.
Bacterial polynucleotide #1450.
US6605709-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADB54303 standard; DNA; 1984 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         . AG.
5.3%;
48.2%;
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                                          (BEHW) BEHRINGWERKE AG.
Query Match
5.3%;
Best Local Similarity 44.3%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                            Local Similarity
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Best Local Similarity
RESULT 1073
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Best Local Similarity
RESULT 1074
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Best Local Similarity
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RESULT 1072
                                                                        Best Local Similarity RESULT 1067
                                                                                                                                                                                                                                                                       12-AUG-2003
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Best Local S
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                                                                                                                                                                                Query Match
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DNA; 13125 BP. cell signalling DNA sequence complementary to#87.
                                                                                                                                                                                                                                                                                                                                                                                                                                         AAS4008 standard; DNA; 9219 BP.
Tumour suppressor gene derived chemically modified sequence #534.
WO200168912-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 4; Length 9219;
                                                                                         Length 6247;
                                                                                                                                                                                                                                                                                                                                                                                            Length 9209;
ABK339923 standard; DNA; 6247 BP.

Human chemically pretreated gene sequence #2 strand
W0200202806-A2.

10-JAN-2002.

[BPIG-) EPIGENOMICS AG.

5.3%; Score 37.6; DB 6; Len
st Local Similarity 45.8%; Pred. No. 82;
                                                                                                                                                                                                                                                                                         LT 1077
ABL34427 standard; DNA; 9209 BP.
Human immune system associated gene SEQ ID NO: 2400.
Human immune system associated gene SEQ ID NO: 2400.
MO20020028-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
EPIG-) ESPIGENOMICS AG.
5.3%; Score 37.6; DB 6; Ler
cer Anatch Similarity 44.0%; Pred. NO. 89;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABLi33032 standard; DNA; 14798 BP.
Human immune system associated gene SEQ ID NO: 1005
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL33227 standard; DNA; 13125 BP.

Human immune system associated gene SEQ ID NO: 1200.

MO200200929-A2.

03-JNA-2002.

(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 6;
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Human metastasis associated gene SEQ ID NO: 110.
WO20017376-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                  ID ABB54279 standard; DNA; 8085 BP.

DE Pretreated genomic DNA region 203.

PN W02003072821-A2.

PD 04-SEP-2003.

PA (EPIG-) EPIGENOMICS AG.

Query Match

Best Local Similarity 47.1%; Pred. No. 87;

RESULT 1077
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Pred. No. 96;
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Pred. No. 98;
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Pred. No. 96;
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Pred. No. 96;
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Pred. No. 93;
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18-0CT-2001.
(EPIG-) EPIGENOMICS AG.
5.3%;
... MAtch
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10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
-- Match -- 17-rity 49.5%;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
6-rv Match 5:3%;
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Best Local Similarity 49.5%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
Query Match 5
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Best Local Similarity
RESULT 1084
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RESULT 1079
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RESULT 1080
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ID ABL34557 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABL70284 standard;
Chemically treated
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
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Best Local Similarity
RESULT 1076
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DNA encoding novel human diagnostic protein #10909 WO200175067-A2.
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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RESULT 1100
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                                                                                    Local Similarity
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                                                   (HYSE-) HYSEQ INC.
                                    11-OCT-2001
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                                                                     Query Match
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                                         03-7AN 2002.
(EPIG-) EPIGENOMICS AG.
Ery Match 5.3%; Score 37.6; DB 6; Length 17528;
                                                                                                                                                                                                                    Length 19787;
                                                                                                                                                                                                                                                                                                                                                    Length 73334;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 73334;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.2%; Score 37.4; DB 5; Length 1767; 58.6%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              77-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 2; Length 361;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 510;
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                                                                                                                                                                                                                                                                                                                                                                           ABL34125 standard; DNA; 73334 BP.
Human immune system associated gene SEQ ID NO: 2098.
WO200200928-A2.
                                                                                                                                               Human immune system associated gene SEQ ID NO: 1423 WO200200928-A2.
             Human immune system associated gene SEQ ID NO: 574 WO200200928-A2.
                                                                                                                                                                                                                                                                                                         03-0302.
03-0302.
(EPIG-) EPIGENOMICS AG.
(ery Match 5.3%; Score 37.6; DB 6;
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human diagnostic protein #609
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5.3%; Score 37.6; DB 6; 51.2%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ry Match
t Local Similarity 48.8%; Pred. No. 54;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 5 28: Score 37.4; DB 5;
                                                                                                                                                                                                                  Score 37.6; DB Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.2%; Score 37.4; 1 62.1%; Pred. No. 50;
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43.3%; Pred. No. 55;
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43.3%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADL44915 standard; DNA; 510 BP.
Human ovarian cancer DNA marker #18805.
WO200170979-A2.
27-SEP-2001.
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ADLJ37966 standard; DNA; 579 BP.
Human ovarian cancer DNA marker #11856.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human ovarian cancer DNA marker #5569.
WO200170979-A2.
 BP.
                                                                                                                                 DNA; 19787 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAZ27709 standard; DNA; 361 BP.
Human DNA marker clone G025.
ABL32601 standard; DNA; 17528
                                                                                                                                                                  WOZUGETO
03-JAN-2002,
(EPIG-) EPIGENOMICS AG.
TWY MATCh Fishisty 47.5%;
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(PROM-) PROMEGA CORP.
                                                                             Ouery Match
Best Local Similarity
RESULT 1085
                                                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 1086
                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 1087
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Best Local Similarity
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DNA encoding novel
WO200175067-A2.
                                                                                                                              ABL33450 standard;
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Best Local Similarity
RESULT 1093
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(HYSE-) HYSEQ INC.
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PN
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ABL70324 standard; DNA; 6181 BP.
Chemically treated cell signalling DNA sequence complementary to#107.
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Signal transduction associated gene modified complementary DNA #105. WO200200926-A2.
                                                                                                                                       Length 2310;
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 Length 1767;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 5852,
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Human immune system associated gene SEQ ID NO: 1749.
                                                                                                                                         DB 12;
                                                                                                                                                                                                                                                                                                                                                                                                                      DB 11;
DB 5;
                                                                                                                                                                                                                                                                              DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    9
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Dictyostellum plasmid Ddp2 containing Rep gene
W09106644-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB
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                                             AD035574 standard; DNA; 2310 BP.
Novel mouse gene sequence #247.
NO2004046310-A2.
O3-JUN-2004.
(FIVE-) FIVE PRIME THERAPEUTICS INC.
Query Match
Eset Local Similarity 60.2%; Pred. No. 74;
5.2%; Score 37.4; 1
58.6%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                      5.2%; Score 37.4; I
62.1%; Pred. No. 77;
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( (EPIG-) EPIGENOMICS AG. 5.2%; Score 37.4; I Guery Match 5.2%; Pred. No. 91;
                                                                                                                                                                                                                                                                              Score 37.4;
Pred. No. 75;
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Pred. No. 90;
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Pred. No. 91;
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Pred. No. 91;
                                                                                                                                                                                                                                                                                                                           ADM01479 standard, cDNA, 2778 BP.
Human cDNA of the invention SEQ ID NO:164.
EP1347046-A1.
24-SEP-2003.
(REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                      AAQ11711 standard; DNA; 2422 BP. Shuttle vector pMUW1530.
WO9106644-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DNA; 5984 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WOSLOCC.
16-MAY-1991.
(UVMA-) UNIV MACQUARIE,
TAY MATCh -- 1 arity 48.7%;
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13-JUN-2002.
(EPIG-) EPIGENOMICS AG.
5.2%;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.2%;
                                                                                                                                                                                                                                        PD 16-MAY-1991.

PA (UYMA-) UNIV MACQUARIE.

Ouery Match

Best Local Similarity 48.7%;

RESULT 1096
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          la-Uc.r.z.v.r.
(EPIG-) EPIGENOMICS AG.
FY Match 5.2%;
'Yoral Similarity 48.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DNA; 6181
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AAS46417 standard; DNA; 6668 BP.
Tumour suppressor gene derived chemically modified sequence #139 WO200168912-A2.
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Human chemically modified disease associated gene SEQ ID NO 179-
WC200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          03-JAN-2002.
(RPIG-) EPIGENOMICS AG. 5.2%; Score 37.4; DB 6; Length 6917; ery Match 5.2%; Pred. No. 93;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match 5.2%; Score 37.4; DB 6; Length 8305; Best Local Similarity 48.4%; Pred. No. 97; RESULT 1111
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 6668;
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                                                   Length 6239;
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                                                                                                                                                                                                                                                                                                                                    Length 6250;
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                                                                                                                                                                                                                                                 ABN80215 standard; DNA; 6250 BP.
Human chemically modified disease associated gene SEQ ID WO200200927-A2.
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Human immune system associated gene SEQ ID NO: 1541.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL33218 standard; DNA; 6668 BP.
Human immune system associated gene SEQ ID NO: 1191.
WC200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AEL34235 standard; DNA; 6917 BP.
Human immune system associated gene SEQ ID NO: 2208.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 6;
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                                                       DB 6;
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                                                             RESULT 1103
TARK28297 standard, DNA, 6239 BP.
DE DNA transcription associated genomic DNA #86.
PN W0200192565-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 37.4; I
Pred. No. 93;
                                                                                                                                                                                             5.2%; Score 37.4; 1
50.3%; Pred. No. 91;
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Best Local Similarity 53.8%; Pred. No. 93;
                                                     Score 37.4;
Pred. No. 91;
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Pred. No. 93;
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Chemically treated apoptosis gene #12.
WC200177164-A2.
18-OCT-2001.
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Novel human cDNA sequence SeqID1.
JP2004097206-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               53.2%;
                                                   5.2%;
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PA (EPIG-) EPIGENOMICS AG.
Query Match
Best Local Similarity 53.8%;
RESULT 1108
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) BPIGENOMICS AG.
WO200200928-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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Query Match
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(EPIG-) EPIGENOMICS AG.
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2002.
2002.
..G-) EPIGENOMICS.
..ery Match
Best Local Similarity 5.
RESULT 1107
ID ARN80162 standa~'
DR Human chemic
PD 03-2020.
PA
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                                                     Query Match
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ID ABL33210
DE Human it
PN WC200200
PD 03-JAN-:
PA (EPIG-)
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AAK73166 standard; DNA; 19965 BP.

Human immune/haematopoietic antigen genomic sequence SEQ ID NO:27978.

Human immune/haematopoietic antigen genomic sequence SEQ ID NO:27978.

90-AUG-2001.

(HUMA.) HUMAN GENOME SCI INC.

1 Score 37.4; DB 4; Length 19965;

ery Match

1 Similarity 47.3%; Pred. No. 1.2e+02;
                                                                                                                                                                                   PN WOLCOLOUS AG.

PD 20-SEP-2001.

PA (EPIG-) EPIGENOMICS AG.

9.24; Score 37.4; DB 4; Length 17144;

Best Local Similarity 53.8%; Pred. No. 1.1e+02;

RESULT 1113

ID ADP72698 standard; DNA; 19080 BP.

DE Renal toxinn progression gene marker #1287.

PD 10-JUN-2004.

PD 10-JUN-2004.

PD (GNE.) GRNE LOGIC INC.

Query Match

S.2%; Score 37.4; DB 12; Length 19080;

Renat Local Similarity 52.2%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match 5.2%; Score 37.4; DB 6; Length 19965;
Best Local Similarity 47.3%; Pred. No. 1.2e+02;
RESULT 1116
ID AAK68939 standard; DNA; 28180 BP.
DE Human immune/haematopoietic antigen genomic sequence SEQ ID NO:23751.
PN WO200157182-A2.
                                                                                                                      AAS46665 standard; DNA; 17144 BP.
Tumour suppressor gene derived chemically modified sequence #387
WO200168912-A2.
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                                                            Length 10591;
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Human secreted protein gene 68 genomic DNA fragment #20
WO200226931-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ID ABL34197 standard; DNA; 37973 BP.

DB Human immune system associated gene SEQ ID NO: 2170.

DP WO200200928-A2.

PD 03-JAN-2002.

PA (EPTG-) EPIGENOMICS AG.

Ouery Match

Best Local Similarity 60.2%; Score 37.4; DB 6; Lem.

RESULT 1118
PD 02-APR-2004.

PA (KAZU-) ZH KAZUSA DNA KENKYUSHO.

PA (PROT-) PROTEIN EXPRESS KK.

Query Match
Best Local Similarity 48.0%; Pred. No. 1e+02;

RESULT 1112
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        04-APR-2002.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 5.2%; Score 37.4; DB 6;
st Local Similarity 47.3%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAD36834 standard; DNA; 59215 BP.

Human transporter protein genomic DNA.

Query Match

5.2*; Score 37.4; DB 8;

Best Local Similarity 70.4*; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 37.4; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5.2%; Score 37.4; DB 4;
47.3%; Pred. No. 1.3e+02;
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Pred. No. 1.6e+02;
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Pred. No. 1.7e+02;
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(HUMA-) HUMAN GENOME SCI INC.
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J.2001.
JMA-) HUMAN GENON.
JUEYY MATCH
Best Local Similarity RESULT 1117
ID ABL34197 stand-
DE Human immur
PN W020020**
PD 03-72**
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ID AAC89888 standard;
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Best Local Similarity
RESULT 1115
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Best Local Similarity
RESULT 1121
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Best Local Similarity
RESULT 1122
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(GEST ) GENSET
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DB 6; Length 692;

Length 887;

DB 3;

Length 843;

5.

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RECULT 1137
ID AAH19205 standard, cDNA, 891 BP.
DE Human secreted protein-encoding gene 7 cDNA clone HPWAY46, SEQ ID NO:46.
PN W0200132910-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human colon cancer antigen encoding cDNA SEQ ID NO:1486 WO200122920-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PD 10-MAY-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Query Match 5.2%; Score 37.2; DB 4;

Best Local Similarity 49.0%; Pred. No. 68;

RESULT 1138
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WO2003038063-A2.
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Human secreted protein encoding sequence #266.
WO200290526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                   5.2%; Score 37.2; DB 49.7%; Pred. No. 67;
                                                                                                                                                                                                  RESULT 1134

ID ABV16074 standard; cDNA; 843 BP.

ID ABV16074 standard; cDNA; 843 BP.

E Human prostate expression marker cDNA 16065.

PN WC200160860-A2.

PD 23-AUG-2001.

PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
5.2%; SCORE 37.2; D'
5.2%; Pred. No. 64;
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08-MAY-2003.

(HUMA-) HUMAN GENOME SCI INC.

5.2%; Score 37.2; D'

5.2%; Pred. No. 68;
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46.8%; Pred. No. 68;
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(HUWA-) HUMAN GENOME SCI INC.
ELY MATCh 5.2%; Score 37.2; I
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49.0%; Pred. No. 68;
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(HUMA-) HUMAN GENOME SCI INC.
ery Match
' """ilarity 49.0%; Pred. No. 68;
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Best Local Similarity 49.0%; Pred. No. 68;
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Gene encoding human secreted protein #445.
W02002102994-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Human secreted protein encoding cDNA WO2002102993-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAC59297 standard; cDNA; 887 BP.
Human secreted protein cDNA #21.
WO200056753-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CDNA; 891 BP
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(HUMA-) HUMAN GENOME SCI INC.
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Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1141 ID ADA56629 standard
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAH34404 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADA40469 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADC73918 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    28-SEP-2000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               05-APR-2001
                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 1136
                                                                  Length 110000;
                                                                                                                                                                 Length 110000,
                                                                                                                                                                                                                                                                                                                                                                              Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 143391;
                                                                                                                                                                                                                                                                           Length 110000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 139308;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 160771;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 299;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 4; Length 540;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 581;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABQ88179 standard; cDNA; 160771 BP.
Human osteoblast differentiation related cDNA SEQ ID NO 86.
W020301-A2.
27-UW-2002.
(GENE-) GENE LOGIC INC.
(PROC ) PROCTER & GAMBLE CO.
ETY MATCh

ETY
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAH35985 standard; cDNA; 540 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:3067.
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADL13648 standard; DNA; 143391 BP.
Osteoarthritis-associated polymorphic nucleotide #180.
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Listeria monocytogenes 4b contig DNA sequence #1086.
11-APR-2002.
                                                          Score 37.4; DB 10;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                  Score 37.4; DB 12;
Pred. No. 1.7e+02;
                                                                                                                                                                 Score 37.4; DB 12;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           vuery Match 5.2%; Score 37.4; DB 10; Best Local Similarity 57.1%; Pred. No. 1.8e+02; RESULT 1128
                                                                                                                                                                                                                                                                        Score 37.4; DB 12;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AD172102 standard; DNA; 299 BP.

Human ovarian cancer DNA marker #4844.

WC200170979-A2.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

ery Match

st Local Similarity 42.9%; Pred. No. 54;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      4,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.2%; Score 37.4; DB 8; 47.3%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 5.28: Score 37.2; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB
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Human cDNA clone (3'-primer) SEQ ID NO:8111.
EP1074617-A2.
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ON-APR-2001.

ON-APR-2001.

ON-APR-2001.

S.2*; Score 37.2; D.

S.2*; Score 37.2; D.

Oney Match

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(HELL-) HELIX RES INST.
5.2%; Score 37.2; I
ory Match 62; Pred. No. 62;
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42.9%; Pred. No. 54;
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RESULT 1130
ID ADL37251 standard; DNA, 299 BP.
DE Human ovarian cancer DNA marker #11141.
PD 27-SEP-2001.
PA (MILL-) MIII,FNNNTHM CHARLES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADB12769 standard; DNA; 139308 BP.
Human PRKR DNA.
DE10128838-A1.
                                                                                                                                                                                                                                                                                                                                                                              5.2%;
                                                       5.2%;
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(INCY-) INCYTE GENOMICS INC.
                                                                                                                                                                 5.2%;
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(GENP-) GENPROFILE AG.
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 1129
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Best Local Similarity
RESULT 1132
                                                                                           Best Local Similarity
                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1124
                                                                                                                                                                                                                                                             Ouery Match
Best Local Similarity
RESULT 1125
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Best Local Similarity
RESULT 1126
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ID ADL13648 standard;
                                                          Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
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ID DE PD PA

Length 891;

Length 891;

DB 4;

Length 891;

Length 891;

DB 10;

Length 891;

DB 8;

Length 891;

10;

DB

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18-OCT-2001
           03-JAN-2002
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                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 1155
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   MEK42900 standard; DNA; 4149 BP.
Genomic sequence #889 encoding novel human connective tissue polypeptide.
WR2010153943.Al.
O2-AUG-2001.
(HUWA-) HUMAN GENOME SCI INC.
ETY MAtch
S.2%; Score 37.2; DB 4; Length 4149;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAK78680 standard; DNA; 4149 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:33492.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 4149;
                                                                                            Arabidopsis thaliana stress regulated gene SEQ ID NO 3866.

Arabidopsis thaliana stress regulated gene SEQ ID NO 3866.

WO200216655-A2.

SCRI ) SCRIPPS RES INST.

A (STRI ) SURGENTA PARTICIPATIONS AG.

Query Match

5.2%; Score 37.2; DB 6; Length 2000;

Best Local Similarity 46.0%; Pred. No. 80;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 8; Length 2695;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 37.2; DB 8; Length 3096;
Pred. No. 88;
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
1ery Match 5.2%; Score 37.2; DB 6; Length 1657;
1ery Match 19.0%; Pred. No. 77;
                                                                                                                                                                                                                                                                                                                            Length 2000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 2602
                                                                                                                                                                                                                                                                                                                                                                          ABX77542 standard; cDNA; 2602 BP.
Differentially expressed breast cancer associated cDNA #37.
US2001156263-A1.
24-0CT-2002.
(CHEN/) CHEN H.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human secretory polynuclectide SPTM SEQ ID NO 127. WC200283876-A2.
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PA (HUMA-) HUMAN GENOME SCI INC.
Query Match 5.2%; Score 37.2; DB 9;
Best Local Similarity 56.6%; Pred. No. 93;
RESULT 1151
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 4;
                                                                                                                                                                                                                                                                                                                            DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADB61146 standard; DNA; 4149 BP.
Connective tissue related genomic DNA #889.
US2003054375-A1.
                                                                                                                                                                                                                                                                                         (SYGN ) SYNGENTA PARTICIPATIONS AG. (SYGN ) SYNGENTA FARTICIPATIONS AG. (EYY Match 5.2%; Score 37.2; I 5.1%; Pred. No. 80;
                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.2%; Score 37.2; 1
51.2%; Pred. No. 85;
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56.6%; Pred. No. 93;
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WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABZ35963 standard; cDNA; 2695 BP
                                                                                                                                                                                                                                            ADA71517 standard; DNA; 2000 BP. Rice gene, SEQ ID 4840. WO2003000898-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 3096 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.2%;
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(INCY-) INCYTE GENOMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
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Best Local Similarity

RESULT 1150

ID ADB61146 stand**

DE Connective '

PN US2003^**

PD 20**
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RESULT 1149
                                               Query Match
Best Local Similarity
RESULT 1143
                                                                                                                                                                                                            Best Local Similarity
RESULT 1144
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Best Local Similarity
RESULT 1148
                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1145
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
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ID ABZ35962
DE Human se
PN WO200283
PD 24-OCT-.7
PA (INCY-)
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DNA; 7145 BP. human genomic DNA #11 associated with DNA adducts
                                                                                                                                                                                  sequence #284.
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(BPIG-) EPIGENOMICS AG.
5.2%; Score 37.2; DB 6; Length 6486;
Alch Match imilarity 46.5%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     6; Length 5763;
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                                     Length 5204;
                                                                                                                                                                                                                                                   03-0CT-2002.
(BPIG-) EPIGENOMICS AG.
ery Match
... similarity 50.6%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 5666;
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RESULT 1156
ID ABQ67049 standard; DNA; 6486 BP.
DE Human anglogenesis associated polynucleotide SEQ ID NO
PN W0200246454-A2.
                                                                                                                                                    ABZ10144 standard; DNA; 5660 BP. Haematopoietic cell proliferation disorder related DNA W0200277272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                          LT 1125.
LT 1125.
LT 1125.
LC 1125.
LC 1125.
Colon cancer analysis related genomic DNA SEQ ID NO:61
W02003101338-A2.
20-FEB-2003.
(EPIG-2003.
(EPIG-) EPIGENOMICS AG.
5.2%; Score 37.2; DB 8; Lengt
sery Match
1.0cal Similarity 50.6%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human immune system associated gene SEQ ID NO: 374.
WO200200928-A2.
03-JAN-2002.
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Query March

BEST Local Similarity 47.1%; Pred. No. 1e+02;

RESULT 1158
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Pred. No. 1e+02;
                                         DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL92234 standard, DNA, 7145 BP. Chemically treated DNA repair gene fragment#22 WO200181622-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human immune system associated gene SEQ ID NO: WO200200928-A2.
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Q.J.AN.-2002.

(EPIG-) EPIGENOMICS AG.

S.2%; Score 37.2; DB (
S.2.%; Score 37.2; DB (
S.2.%; Score 37.2; DB (
S.2.%; Store 37.2; DB (
S.2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.2%; Score 37.2; DB 56.6%; Pred. No. 1e+02;
PA (EPIG-) EPIGENOMICS AG.
Query Match
Best Local Similarity 45.6%; Pred. No. 98;
RESULT 1152
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity 50.64; Pred. RESULT 1154

ID ABL32182 standard; DNA; 5763 BP. DE Human immune system associated gen. PN W20020020928-A2. PD 03-GAN-2002. PA (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (ORIG-) ORIGENE TECHNOLOGIES INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAL60047 standard; cDNA; 7096 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL32401 standard; DNA; 7167 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DNA; 6145 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity 47.1%;
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Chemically treated
WO200177378-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best_Local Similarity RESULT 1157
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Best Local Similarity
RESULT 1153
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AAS46787 standard; DNA; 61020 BP.
Tumour suppressor gene derived chemically modified sequence #513
WO200168912-A2.
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DE Prostate tumour related genomic DNA sample #9.

DE Prostate tumour related genomic DNA sample #9.

PN M020013044-A2.

PA (EPIG-) EPIGENOMICS AG.

Query March

Best Local Similarity 55.4%; Pred. No. 1.3e+02;

DE Human renal/prostate carcinoma associated DNA SEQ ID NO.17.

PN W02002103041-A2.

PN W02002103041-A2.

PA (EPIG-) EPIGENOMICS AG.

Query March

S.2%; Score 37.2; DB 8; Length 18997;

PN W02002103041-A2.

PN W02002103041-A2.

PA (EPIG-) EPIGENOMICS AG.

Query March

S.2%; Score 37.2; DB 8; Length 18997;

PA (EPIG-) EPIGENOMICS AG.

S.2%; Score 37.2; DB 8; Length 18997;

DE Human renal/prostate carcinoma associated DNA SEQ ID NO.17.

PA (EPIG-) EPIGENOMICS AG.

S.2%; Score 37.2; DB 8; Length 18997;

DE Human renal/prostate carcinoma associated DNA SEQ ID NO.17.

PA (EPIG-) EPIGENOMICS AG.

Ouery March

Best Local Similarity 55.4%; Pred. No. 1.3e+02;
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              Length 18997;
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Mouse Nfkbl carcinoma associated gene, SEQ ID NO:1127
WO2003057146-A2.
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Pred. No. 1.8e+02;
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Pred. No. 1.8e+02;
           5.2%; Score 37.2; DB 6; 55.4%; Pred. No. 1.3e+02;
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(EPIG-2001.
(EPIG-2001.
(EPIG-2001.
(EPIG-2001.) EPIGENOMICS AG.
5.2%; Score 37.2; DB 4;
(ery Match 51.2%; Pred. No. 1.6e+02;
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Pred. No. 1.8e+02;
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Pred. No. 1.9e+02;
                                                                                                                                                                                                                                         Score 37.2; DB 6;
Pred. No. 1.3e+02;
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Pred. No. 1.9e+02;
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Pred. No. 1.8e+02;
        Query Match
Best Local Similarity 55.4%; Pred. No. 1.3
RESULT 1170
ID ABK33948 standard; DNA; 18997 BP.
DE Human DNA for staging of Astrocytomas #16.
PN WO200202808-A2.
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WO2003080808-A2.
02-OCT-2003.
(SAGR-) SAGRES DISCOVERY.
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WO2003039484-A2.
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PA (BPIG-) EPIGENOMICS AG.
Query Match
Best Local Similarity 55.4%;
RESULT 1171
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15-MAY-2003.
15-MAY-2003.
15-MAY-201.
5.2*;
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Local Similarity 65.9%;
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(SAGR-) SAGRES DISCOVERY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             17-JUL-2003.
(SAGR-) SAGRES DISCOVERY.
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RESULT 1173
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Mouse Nfkb1 gene.
WO2003008583-A2.
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Best Local Similarity
RESULT 1178
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Best Local Similarity
RESULT 1180
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Signal transduction associated gene modified complementary DNA #37.
W0200200926-A2.
W03-0AN-2002.
(BPIGENOMICS AG.
5.2%; Score 37.2; DB 6; Length 13133;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAS46699 standard; DNA; 8711 BP.
Tumour suppressor gene derived chemically modified sequence #422 WO200168912-A2.
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Tumour suppressor gene derived chemically modified sequence #495.
WO20168912-A2.
20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                         ADE84178 standard; DNA; 8666 BP.
Human lymphoid cell proliferative disorder gene derived DNA #114
WO2003044226-A2.
                                                                                                                                                                                                                                                                 Length 8666;
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                             Length 7167;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length
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Streptococcus pneumoniae genome fragment SEQ ID NO:17
WO9818931-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human immune system associated gene SEQ ID NO: 1874 WO200200928-A2.
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Human immune system associated gene SEQ ID NO: 254
WO200200928-A2.
03-JAN-2002.
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Human immune system associated gene SEQ ID NO: 543.
                                                                                                                                                                                                                                                     5.2%; Score 37.2; DB 10; 50.6%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                 5.2%; Score 37.2; DB 6; 51.9%; Pred. No. le+02;
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(HUWAL) HUMAN GENOME SCI INC.
(ETY Match 5.2%; Score 37.2; DB 2;
ery Match 7.00 1.1e+02;
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(EPIG-) EPIGENOMICS AG.
2ry Match 5.2%; Score 37.2; DB 6;
-- rann Similarity 45.7%; Pred. No. 1.2e+02;
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03-JAN-2002.

(BPC) EPIGENOMICS AG.

(ERY Match

5.2%; Score 37.2; DB 6;

6ry Match

7.2e+02;
                                                                                                           ADB54240 standard; DNA; 8666 BP. Pretreated genomic DNA region 164 WO2003072821-A2.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1162
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Best Local Similarity
RESULT 1163
                                            Best Local Similarity
RESULT 1161
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RESULT 1164
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Best Local Similarity
RESULT 1165
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Best Local Similarity
RESULT 1166
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RESULT 1168
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RESULT 1169
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                                Query Match
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Human liver single exon probe, SEQ ID No 8690. WO200157273-A2.

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AAI39617 standard; DNA; 599 BP.
Probe #8303 used to measure gene expression in human placenta sample.
WC200157272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABSO8446 standard; DNA; 453 BP.
Human genome-derived single exon probe from lung SEQ ID No 8437.
WO200186003-A2.
                                                                                                Length 113515;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 1187
ID AAKSJS92 standard; DNA; 599 BP.
DE Human bone marrow expressed single exon probe SEQ ID NO: 8449.
PN WO200157276-A2.
                                                                                                                                                ID AAK33570 standard; DNA; 453 BP.

ID Human bone marrow expressed single exon probe SEQ ID NO: 8127.

DE Human bone marrow expressed single exon probe SEQ ID NO: 8127.

PN W0200157276-A2.

PD 09-AUG-2001.

PA (MOLE-) MOLECULAR DYNAMICS INC.

QUERY Match

Bost Local Similarity 52.2%; Pred. No. 65;

RESULT 1182
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69;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 4; Length 599 69;
                                                                                                                                                                                                                                                                                              AAK07727 standard; DNA; 453 BP.
Human brain expressed single exon probe SEQ ID NO: 7718.
WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human brain expressed single exon probe SEQ ID NO: 8011 WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABASS751 standard; DNA; 599 BP.
Human foetal liver single exon nucleic acid probe #8056
WC200157277-A2.
ABL34174 standard; DNA; 113515 BP.
Human immune system associated gene SEQ ID NO: 2147.
WO200200928-A2.
                                                                                                  Score 37.2; DB 6;
Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 4;
69;
                                                                                                                                                                                                                                                                                                                                                                                     DB 4;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human liver single exon probe, SEQ ID No 8348 WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB /
65;
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69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Les exon probe 1

"wery Match 5.2%; Score 37; DB
Best Local Similarity 52.2%; Pred. No. 65;
RESULT 1185
ID ABA59751 standard; DNA; 599 HP
PN WC20015727-A2
PD 09-AUG-777-A2
PA ...
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(MOLE-) MOLECULAR DYNAMICS INC.
ery Match
5.2%; Score 37;
                                                                                                                                                                                                                                                                                                                                          09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(ery Match
15.2%; Score 37;
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ry Match
t Local Similarity 52.2%; Pred. No.
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Query Match
Best Local Similarity 52.2%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
5.2%; Score 37;
lery Match 52.2%; Pred. No.
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(MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABS33700 standard; DNA; 599 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                           ABS33358 standard; DNA; 453 BP.
                                                                  03-JAN-2004.
(EPIG-) EPIGENOMICS AG.
FY MACCh 5-28;
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..2001.
..LE-) MOLECULAR L
..LE-Y MATCH
Best Local Similarity ..
RESULT 1188
ID AAK08020 stand>"
DE Human brair
PN WO2001="
PD 09-"
                                                                                                                 Best Local Similarity RESULT 1181
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Best Local Similarity
RESULT 1184
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Best Local Similarity
RESULT 1189
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RESULT 1186
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Best Local Similarity
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                                                                                                    Query Match
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ACC69470 standard; cDNA; 3078 BP.
Human malignant neoplasm related protein encoding cDNA SEQ ID NO:11.
W02003025135-A2.
                                                                                Oligonuclectide for detecting cytosine methylation SEQ ID NO 29110. WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                              PD 07-FEB-2001.

PA (HELI-) HELIX RES INST.

Query Match
5.2%; Score 37; DB 4; Length 1385;
Best Local Similarity 49.8%; Pred. No. 82;
RESULT 1193
ID ABS76410 standard; CDNA; 1385 BP.
DE CDNA encoding human ovarian cancer marker OV28.

PN WO2021928-A2.
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                               Length 599;
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Chemically treated cell signalling DNA sequence#120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAF83292 standard; cDNA; 2110 BP.
P. rhizinflata eglA protein encoding partial cDNA.
US6222028-B1.
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94;
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97;
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PA (EPIG-) EPIGENOMICS AG.

QUETY MATCh

Best Local Similarity 46.4%; Pred. No. 77;

RESULT 1191.
                                                                                                                                                                                                                                                                              07-MAR-2002.

07-MAR-2002.

(EPIG-) EPIGENOMICS AG.

5.2%; Score 37; DB 6;

lery Match

5.2%; Pred. No. 77;
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82;
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PA (BOYC-) BOYCE THOMPSON INST PLANT RES.
Query Match 5.2%; Score 37; DB 2;
Best Local Similarity 50.7%; Pred. No. 1e+02;
RESULT 1198
                                   4
                                 DB
69;
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PD 09-AUG-2001.

PA (MOLE-) MOLECULAR DYNAMICS INC.

Query Match 5.2%; Score 37;

Best Local Similarity 52.2%; Pred. No.

RESULT 1190
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Pred. No.
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50.2%; Pred. No.
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TD AAH1775 standard; cDNA, 2545 BP.
DE Human cDNA sequence SEQ ID NO:17414.
                                                                                                                                                                                                                                                                                                                                                              Human cDNA sequence SEQ ID NO:14774.
EP1074617-A2.
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Banana ripening fruit Gluc. DNA.
WO9915668-A2.
                                                                                                                                                                                                                                 DNA; 1030 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             19-SEP-2002.
(MILL-) MILLENNIUM PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PD 27-MAR-2003.

PA (GENE-) GENE LOGIC INC.

QUETY MATCh

Best Local Similarity 53.9%;

RESULT 1197
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(SINI-) ACAD SINICA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Local Similarity
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Best Local Similarity
RESULT 1192
                                                                                                                                                                                                                                    ABQ42519 standard;
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ID ACC69470 standard;
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Length 6602;

WO200202807-A2.

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Signal transduction associated gene modified complementary DNA #82.

Signal transduction associated gene modified complementary DNA #82.

NOOJOSO0256-A2.

OBJURN SOOR

A (BPIG-) EDIGENOMICS AG.

S.2%; Score 37; DB 6; Length 11907;

Best Local Similarity 50.3%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAS46355 standard, DNA, 14537 BP.
Tumour suppressor gene derived chemically modified sequence #77.
WO200168912-A2.
                                                                                                                                                                                                                                                                       Length 10160;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 10323;
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                                                                                                                                                                                                                                                                                                                                                                                         Query Match 5.2%; Score 37; DB 6; Length 8346; Beet Local Similarity 56.6%; Pred. No. 1.2e+02; RESULT 1210
                                                                                                                                                                                                                                              Length 6729;
                                                                                                                                                           Human angiogenesis associated polynucleotide SEQ ID NO W0200246454-A2.
13.-JUN-2002.
(EPIG-) EPIGENOMICS AG.
5.2%; Score 37; DB 6; Length 6 et Local Similarity 48.4%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            איאי DNA; 10323 BP.
Plant transformation binary vector, plasmid pDAB1542.
W09950430-A2.
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Chemically pretreated metabolism associated gene #8
WO200176451-A2.
                                                                                  Score 37; DB 10;
Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5.2%; Score 37; DB 2; 53.0%; Pred. No. 1.3e+02;
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(EPIG-) EPIGENOMICS AG.
sry Match 5.2%; Score 37; DB 6; 1
... Tarnal Similarity 46.6%; Pred. No. 1.3e+02;
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Pred. No. 1.3e+02;
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53.0%; Pred. No. 1.3e+02;
07-AUG-2003.
(PLAN-) PLANET BIOTECHNOLOGY INC.
(LARR) LARRICK J W.
(WYCO/) WYCOFF K L.
                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAV63723 standard; DNA; 10160 BP.
Vector plasmid pDAB406.
WO9856921-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAV63724 standard; DNA; 11784 BP.
Vector plasmid pDAB411.
WO9856921-A1.
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Vector plasmid pDAB419.
WO9856921-A1.
                                                                                                                                        ABQ67153 standard; DNA; 6729 BP.
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(DOWC ) DOW AGROSCIENCES LLC.
                                                                                5.2%;
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(DOWC ) DOW AGROSCIENCES LLC.
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(DOWC ) DOW AGROSCIENCES LLC.
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Best Local Similarity 53.0%;
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                                                                              Query Match
Best Local Similarity
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Best Local Similarity
RESULT 1213
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                                                                                                                                                                                                                                                                               RESULT 1209
ID ABK28328 standard;
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Best Local Similarity
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DNA derived from between T-DNA borders of plasmid pGPTV-kan-ocs-ATR-1gA2.
WO2003064992-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL70372 standard; DNA; 6070 BP.
Chemically treated cell signalling DNA sequence complementary to#131.
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAS46663 standard; DNA; 6076 BP.
Tumour suppressor gene derived chemically modified sequence #385.
WO200168912-A2.
                                                       Score 37; DB 6; Length 5504;
Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                            Score 37; DB 6; Length 6070; Pred. No. 1.1e+02;
                                                                                                                                                                                                                   Length 5504;
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PA (EPIG-) EDIGENOMICS AG.
Query Match
5.2%; Score 37; DB 6; Length 6290;
Best Local Similarity 54.9%; Pred. No. 1.1e+02;
                                                                                                 AAS6130B standard; DNA; 5504 BP.
Human gene regulation-associated gene oligonucleotide #263.
WO200177375-A2.
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Human angiogenesis associated polynucleotide SEQ ID NO.
WO200246454-A2.
                                                                                                                                                                                                                                                                     ABL33679 standard; DNA; 6070 BP.
Human immune system associated gene SEQ ID NO: 1652
WO200200928-A2.
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Human immune system associated gene SEQ ID NO: 2192
WO200200928-A2.
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Human immune system associated gene SEQ ID NO: 1019
WO200200928-A2.
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18-00-12001.
(EPUG-) EPIGENOMICS AG.
5.2%; Score 37; DB 6; Lerv Match Firsty 51.7%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                ABL34579 standard; DNA; 6070 BP.
Human metastasis associated gene SEQ ID NO: 132.
WO200177376-A2.
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Pred. No. 1.1e+02;
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Pred. No. 1.1e+02;
  MOZOVIII
(EPIG) EPIGENOMICS AG.
(arv Match '-:'arity 51.7%;
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18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
TYV MATCH --1arity 52.2%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                    Best Local Similarity
RESULT 1199
                                                                                                                                                                                                                                 Best Local Similarity
RESULT 1200
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Best Local Similarity
RESULT 1201
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RESULT 1202
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RESULT 1204
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(EPIG-) EPIGENOMICS AG.
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                                                                                                                                                                                                                                                                                                                                              Query Match
                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAS45323 standard; DNA; 17848 BP.
Chemically pretreated complementary DNA associated with cell cycle #14.
WO200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                 ABL70608 standard; DNA; 15518 BP. Chemically treated cell signalling DNA sequence complementary to#249.
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           cuery Match 5.2%; Score 37; DB 4; Length 17848; BESTL 1.20cal Similarity 44.0%; Pred. No. 1.46+02; DE Human chemically pretreated company w0200202806-A2.

PN W020028806-A2.

PN W020028806-A2.
PD 20-SEP-2001.
PA (BPIG-) EPIGENOMICS AG.
Query Match 5.2%; Score 37; DB 4; Length 14537;
Best Local Similarity 45.2%; Pred. No. 1.3e+02;
RESULT 1217
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 17848;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.2%; Score 37; DB 3; Length 17111; 53.0%; Pred. No. 1.4e+02;
                                                                                                                                                                                   Length 15518;
                                                                                                                                                                                                                                                                                                                 Length 15518;
                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 37; DB 6; Length 15518; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 15676;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABK28164 standard; DNA; 17848 BP.
DNA transcription associated complementary genomic DNA #19.
WO200192565-A2.
06-DEC-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAD01289 standard; DNA; 17111 BP.
Arabidopsis transformation binary vector, pArAct2Af-bin.
W0200032800-Al.
(8-JUN-2000.
(DOWC ) DOW AGROSCIENCES LLC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAD01290 standard; DNA; 17116 BP.
Arabidopsis transformation binary vector, pAfAct2Af-bin.
WO200032800-A1.
                                                                                           ABL34173 standard; DNA; 15518 BP.
Human immune system associated gene SEQ ID NO: 2146.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAD01288 standard; DNA; 15676 BP.
Arabidopsis transformation binary vector, pAct2-bin
W0200032800-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           PD 08-JUN-2000.
PA (DOWC) DOW AGROSCIENCES LLC.
Query Match 5.2%; Score 37; DB 3; I Best Local Similarity 53.0%; Pred. No. 1.4e+02; RESULT 1223
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PA (DOWC) DOW AGROSCIENCES LLC.
Query Match
Best Local Similarity 53.0%; Pred. No. 1.4e+02;
RESULT 1221
                                                                                                                                                                                   Score 37; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                  Human metastasis associated gene SEQ ID NO: 178
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 37; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                 Score 37; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                     ABL34625 standard; DNA; 15518 BP.
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10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.2%;
                                                                                                                                                                                                                                                                                    18-OCT-20U1.
(EDIG-) EPIGENOMICS AG.
QUETY MATCH 5.2%;
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ry Match 5.2%;
t Local Similarity 45.6%;
                                                                                                                                                                                   5.2%;
                                                                                                                                                     03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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...d.) EPIGENOMICS
...ery Match
Best Local Similarity
RESULT 1225
ID ABKZ8164 stand
DE DNA trand
PN WO200
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RESULT 1220
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                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
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AANSO182 standard; DNA; 24596 BP. Complete nucleotide sequence of the T-DNA region of the octopine Ti plasmid pTi15955.
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Bovine EST associated with lactation/muscle/fat deposition #1221.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAS46745 standard; DNA; 38342 BP.
Tumour suppressor gene derived chemically modified sequence #469.
WO200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 29993;
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Length 17848;
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                                                                                                                                                                                                                                                                                                                                                                                             Length 24596;
                                                                                                                                                                                      Length 24593;
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WO200200926-A2.
03-JAN-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADB37660 standard; DNA; 29993 BP.
Human chemically pretreated EYA4 gene SEQ ID NO:2.
WO2003072812-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              19-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AALU7623 standard; cDNA; 391 BP.
Human breast cancer expressed polynucleotide 80.
W0200151628-A2.
                                                                                                                                            PA (AGR. AGRIGENETICS RES ASSOC LTD.

PA (LUBR) LUBRIZOL GENETICS INC.

Query March

Bost Local Similarity 53.0%; Pred. No. 1.5e+02;

RESULT 122.
                                                                                                                                                                                                                                                                                                                                                                                               5.2%; Score 37; DB 1; 1
53.0%; Pred. No. 1.5e+02;
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    5.2%; Score 37; DB 6; I
44.0%; Pred. No. 1.4e+02;
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Pred. No. 1.7e+02;
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Best Local Similarity 50.9%; Pred. No. 1.66+02;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
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49.4%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.2%; Score 36.8; I
52.6%; Pred. No. 69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.2%; Score 36.8; 358.0%; Pred. No. 71;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human ovarian cancer DNA marker #18240. #0200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAI81705 standard; cDNA; 403 BP.
Human polynucleotide SEQ ID NO 1765.
WO200164835-A2.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                       19-JUN-1985.
(AGRK) AGRIGENETICS RES ASSOC LTD.
(LUBR) LUBRIZOL GENETICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABK31506 standard; DNA; 38342 BP.
                                                              AAN50226 standard; DNA; 24593 BP.
                                                                                    Sequence of opine synthase gene. EP140556-A. 08-MAY-1985.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  WOACOLL
(B-DG-) EPIGENOMICS AG.
(B-DG-) EPIGENOMICS AG.
(B-DG-) EPIGENOMICS AG.
(B-DG-) AGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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RESULT 1228
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                   Best Local Similarity RESULT 1226
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ADJ75071 standard; DNA; 1331 BP.
Marker gene SEQ ID NO:323.
EP1394274-A2.
                                                     03-MAR-2004.
(GENO-) GENOX RES INC.
                                                                                                          Best Local Similarity
                                                                                                                                                                                                                                           Best Local Similarity RESULT 1244
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
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RESULT 1246
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(HMGE-) HMGENE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL92207 standard, Chemically treated WO200181622-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
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Best Local Similarity
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                                                                                           Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAA02542 standard; cDNA; 1199 BP.
Human colon cancer cell line polynucleotide sequence SEQ ID NO:2533.
W09958675-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABX46983 standard; cDNA; 453 BP.
Bovine EST associated with lactation/muscle/fat deposition #12148.
US2002137139-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 1199;
                                                                                                        Length 408;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 507;
                                                                                                                                                                                                                                                                                                              Length 416
                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 6; Length 453
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 8; Length 453;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 567
                                                                                                                                                                                                                                                                                                                                                                        AmEPV apoptosis associated protein (AMV193) encoding WO200212526-A2.
                                                                                                      DB 8;
                                                                                                                                                                                                                                                                                                          5.2%; Score 36.8; DB 9; 32.7%; Pred. No. 71;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
5.2%; Score 36.8; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 5.2%; Score 36.8; DB 5;
er Local Similarity 56.7%; Pred. No. 76;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 3;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABZ37586 standard; DNA; 1269 BP.
Candida albicans essential gene SEQ ID NO 6873.
WO200253728-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABV57839 standard; cDNA; 507 BP.
Human prostate expression marker cDNA 57830.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABV58219 standard; cDNA; 567 BP.
Human prostate expression marker cDNA 58210.
WO200160860-A2.
                                                                                                  5.2%; Score 36.8; I 53.5%; Pred. No. 71;
                                                                                                                                                                                                                                                                                                                                                                                                                                             5.2%; Score 36.8; I 50.6%; Pred. No. 73;
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                                                                                                               Best Local Similarity 53.5%; Prec RESULT 1235
ID ACH27023 standard; cDNA; 416 BP. DE Human adult covary cDNA #5403.
PN US2003073623-A1.
                                                                                                                                                                                                                                                                                                                                                          ABL56232 standard; DNA; 453 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.2%;
                                                                                                                                                                                                                (DRMA) DRYANAC R T.
(LABA/) LABAT I.
(STAC/) STACHE-CRAIN B.
(DICK/) DICKSON M C.
(JONE/) JONES L W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11-JUL-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         26-SEP-2002.
(BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                 MATHIALAGAN N.
                                                                                                                                                                                                                                                                                                                                                                                                          14-FEB-2002.
(UYFL ) UNIV FLORIDA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 1241
ID AB222586 standard; DN
DE Candida albicans esse
PN W0200253728-A2.
PD 11-UUL-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (CHIR ) CHIRON CORP. (HYSE-) HYSEQ INC.
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Best Local Similarity
RESULT 1240
                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1236
                                                                    (TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1237
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
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RESULT 1239
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RESULT 1242
               26-SEP-2002.
(BYAT/) BYATT J C.
(MATH/) MATHIALAGA
US2002137139-A1.
                                                                                                                                                                                                     17-APR-2003
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                                                                                                  Query Match
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                                                                 (TAON/)
                                                                                                                                                  ID
DE
PN
PA
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DB 12; Length 1331;
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DNA repair gene fragment complementary to#8
                                                                                                                                                      Length 1387;
                                                                                                                                                                                                                                                                                                                                           DB 6; Length 1387;
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(EPIG-) EPIGENOMICS AG.
(EY/ Match 5.2%; Score 36.8; DB 6; Length 5826;
(er/ Match 5.2%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADM67057 standard, DNA; 2188 BP.
Human homologue of murine adipocyte specific DNA SeqID 192
WO2004011618-A2.
                              RESULT 1243

ID AAS29132 standard; cDNA; 1387 BP.

DE CDNA encoding for human DNA-binding protein #103

PN WO200155162-A1.
                                                                                                                                                      5
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  / Match 5.2%; Score 36.8; DB 3; Local Similarity 46.1%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                RESULT 1245
ID ADC25266 standard; cDNA; 1387 BP.
DE Human cDNA from extracellular matrix gene 103.
PP US2003409650-A1.
PP USA02409650-A1.
PP INC.
F. Score 36.8; DB
                                                                                                                                                                                                  ABS68272 standard; cDNA; 1387 BP.
cDNA encoding human DNA-binding protein #103.
US2002102638-A1.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(RUBE/) BARASH S C.
                                                                                                                                                    DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               RESULT 1248
ID AAF22346 standard; cDNA; 2058 BP.
DE Human secreted protein gene 31 SEQ ID NO:41.
PN W020061748-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     O3-JUN-2004.

(FIVE-) FIVE PRIME THERAPEUTICS INC.

ery Match

5.2%; Score 36.8; I
     Score 36.8; 1
Pred. No. 91;
                                                                                                                                                5.2%; Score 36.8; I
59.6%; Pred. No. 92;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.2%; Score 36.8; I 59.6%; Pred. No. 92;
                                                                                                                                                                                                                                                                                                                                       5.2%; Score 36.8; 59.6%; Pred. No. 92;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 36.8; I
Pred. No. 99;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADB58887 standard; DNA; 1967 BP.
Toxicity-related gene, SEQ ID 3913.
WO2003064624-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADO35573 standard; DNA; 1520 BP.
Novel mouse gene sequence #246.
WO2004046310-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WO2003000-0...
07-AUG-2003.
(GENE-) GENE LOGIC INC.
5.2%; SC
                                                                                                         02-AUG-2001,
(HUMA-) HUMAN GENOME SCI INC.
5.2%;
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(HUMA-) HUMAN GENOME SCI INC.
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02-JAN-2003
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                                                                                                                              Query Match
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                                                                                                                                                       Duery Match
Best Local Similarity 50.0%; Pred. No. 1.3e+02;
Best Local Similarity 50.0%; Pred. No. 1.3e+02;
RESULT 1252
RESULT 1252
DE ABL70386 standard; DNA; 7321 BP.
DE Chemically treated cell signalling DNA sequence complementary to#138.
PN W0200202807-A2.
PN W0200202807-A2.
PN WG200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABZ10246 standard; DNA; 8056 BP.
Haematopoietic cell proliferation disorder related DNA sequence #386.
WO200277272-A2.
                 ABX31425 standard; DNA; 7321 BP.
Signal transduction associated gene modified complementary DNA #134.
WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 36.8; DB 8; Length 8056; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 7346;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 7461;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 7749;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 7321
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAS61394 standard; DNA; 7341 BP.
Human gene regulation-associated gene oligonucleotide #349.
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Human gene regulation-associated gene oligonucleotide #293 WC200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL33784 standard; DNA; 7461 BP.
Human immune system associated gene SEQ ID NO: 1757.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL70435 standard; DNA; 7749 BP.
Chemically treated cell signalling DNA sequence#163
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL32345 standard; DNA; 7346 BP.

Human immune system associated gene SEQ ID NO: 318.

Human immune system associated gene SEQ ID NO: 318.

103-JAN-2002.

103-JAN-2002.

103-JAN-2002.

104-JALCH

105-JAN-2002.

105-J
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(EPIG-) EPIGENOMICS AG.

5.2%; Score 36.8; DB 6;

10-27 Match 5.2%; Pred. No. 1.3e+02;
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03-OCT-2002.
(BPIG-) EPIGENOMICS AG.
5.2%; Score 36.8; DB 8; Lery Match
7.1%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.2%; Score 36.8; DB 6; 51.2%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                    5.2%; Score 36.8; DB 6; 50.0%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 36.8; DB 6;
Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAS61338 standard; DNA; 7321 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WOZUCZ-2002.
03-007-2002.
(EPIG-) EPIGENOMICS AG.
5.2%;
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18-001-2001.
(EPIG-) EPIGENOMICS AG.
5.2%;
                                                                                           WOLVOLL
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5-2%;
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-2002.

-2002.

-4ery March

Best Local Similarity A

RESULT 1259

ID ABZ10100 standa-
DE Haematopoia*
PN W020027

PA 03-
PA
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2002.
1G-) EPIGENOMIC.
L-GT MATCH
Best Local Similarity
RESULT 1256
ID ABL33784 stando
DE Human immur
PN W02002"
PD 03
                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 1255
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Best Local Similarity
RESULT 1258
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Best Local Similarity
RESULT 1260
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RESULT 1251
1D ABX3142
DE Signal
PN WO20020
PD 03-JAN-
PA (EPIG-)
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ABL70136 standard; DNA; 11416 BP.
Chemically treated cell signalling DNA sequence complementary to#13.
WO200202807-A2.
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ABL49302 standard; DNA; 10467 BP.
Human polymucleotide associated with DNA replication SEQ ID NO
WO200177377-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 19124;
                                                        Length 19124;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 11416;
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                                                                                                                                                                                                                                                  Length 11416;
                                                                                                                                                                                                                                                                                                                                                                                                      Length 11416,
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Human gene regulation-associated gene oligonucleotide #19
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.

PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

Query Match
5.2%; Score 36.8; DB 10;
Best Local Similarity 51.2%; Pred. No. 1.8e+02;

RESULT 1268
                                                                                                                                                                            92
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RESULT 1267
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PA (USSH) US DEPT HEALTH & HUMAN SERVICES.

Query Match
Best Local Similarity 45.5%; Pred. No. 1.6e+02;

RESULT 1265
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Ouery Match 5.2%; Score 36.8; DB 3; Best Local Similarity 45.5%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 36.8; DB 6;
Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                        PD 10-JAN-2002.

PA (EPIG-) EPIGENOMICS AG.

Query Match

Guery Match

Best Local Similarity 47.4%; Pred. No. 1.4e+02;

RESULT 1263
                                                                                                                                                                                                                                                  Score 36.8; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 36.8; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                      Human immune system associated gene SEQ ID NO: W0200200928-A2.
03-JAN-2002.
(EPIGENOMICS AG. 5.2%; Score 36.8; DB 6; SYY Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAZ98287 standard; DNA; 19124 BP.
Plasmodium var-7 polypeptide encoding DNA.
US5993827-A.
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(USSH ) US DEPT HEALTH & HUMAN SERVICES.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL56202 standard; DNA; 50000 BP.
AmEPV genome fragment#4.
WO200212526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADC87242 standard, DNA; 38918 BP.
Human GPCR gene SEQ ID NO:1695.
EP1270724-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 32176 BP
                                                                                                                                                            DNA; 11416 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5.2%;
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                                                                                                                                                                                                                                                      5.28;
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Plasmodium var-7 gene.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      14-FEB-2002.
(UYFL ) UNIV FLORIDA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 1269
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RESULT 1264
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ID AAL05628 standard;
                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 1262
                                                                                                                  Best Local Similarity RESULT 1261
                                                                                                                                                            ABL32119 standard;
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ABX40612 standard; cDNA; 241 BP.
Bovine EST associated with lactation/muscle/fat deposition #5777.
US2002137139-A1.
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          5.1%; Score 36.6; DB 6; Length 200; 62.6%; Pred. No. 68;
                                                                                                                                                                                                                                            DB 8; Length 241;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 456;
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                                                                                                                                                                                                                                                                                                                                                                                                                Length 392;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH48299 standard; cDNA; 2863 BP.
Human phosphoenol pyruvate carboxylase 81 coding sequence.
WO200155412-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADI35832 standard; DNA; 4018 BP.
Mouse proopionmelanocortin (POMC) regulatory region
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.1%; Score 36.6; DB 10; 49.2%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                              5,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5.1%; Score 36.6; DB 4; 49.2%; Pred. No. 81;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (BIOD-) BIODOOR GENE TECHNOLOGY LTD SHANGHAI.

ry Match 5.1%; Score 36.6; DB 4;

t Local Similarity 49.8%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        5.1%; Score 36.6; DB 3; 49.2%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                 LT 1280
ADL4353 standard; DNA; 392 BP.
ADL4353 standard; DNA marker #17543.
WC200170979-A2.
27-SEP-2001.
(Mill-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.1%; Score 36.6; DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human prostate expression marker cDNA 52755 WO200160860-A2.
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WO200061625-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                          5.1%; Score 36.6; I
50.9%; Pred. No. 71;
                                                                                                                                                                                                                                                                                                                                                                                                              5.1%; Score 36.6; I
41.1%; Pred. No. 78;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.1%; Score 36.6; D
58.9%; Pred. No. 84;
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48.8%; Pred. No. 81;
                                                                                                                                                                                                                                                                                                                                                                                                                                                 RESULT 1281

D. AA183622 standard; cDNA; 456 BP.
DE Human polymucleotide SEQ ID NO 3682.
PN WO200164835-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADF74262 standard; DNA; 1239 BP.
Enteroaggregative E_coli aatA DNA.
US2003180315-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAC93445 standard; cDNA; 2904 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CDNA; 456
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  CDNA; 559
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (HUMA-) HUMAN GENOME SCI INC (ROSE/) ROSEN C A.
                                                                                                                                                                             z
                                                                                                                            26-SEP-2002.
(BYAT/) BYATT J C.
(MATH) MATHIALAGAN N
(TAON/) TAO N.
(WARR/) WARREN W C.
Query Match
Best Local Similarity
RESULT 1279
                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
                                                                                                                                                                                                                                                         Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                07-SEP-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
RESULT 1282
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABV56478 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABV52764 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             25-SEP-2003.
(NATA/) NATARO J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      19-OCT-2000
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                                                                                                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                             AAK86026 standard; DNA; 54877 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:40838.
WO200157182-A2.
                 Human immune/haematopoietic antigen genomic sequence SEQ ID NO:40837 WO200157182-A2
                                                                                                                                                                                                                                                                                                                               ABZ74225 standard; DNA; 54877 BP.
Secreted protein gene 215 genomic fragment HNGAM58, SEQ ID NO:1372.
WO200277013-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.2%; Score 36.8; DB 12; Length 110000; 45.3%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5.2%; Score 36.8; DB 12; Length 110000; 45.3%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.2%; Score 36.8; DB 10; Length 54877; 50.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                        03-0CT-2002.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
5.2%; Score 36.8; DB B; Length 54877;
...* Aimilarity 50.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 110000;
                                                    09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
- 5.2%; Score 36.8; DB 4; Length 54863;
- 7.23 Similarity 50.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                             Length 54877;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 54877;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human secreted protein encoding genomic DNA SEQ ID NO 1314 WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL85806 standard; cDNA; 200 BP.
Human ovarian cancer related cDNA clone SEQ ID NO:8784.
WO200192581-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.2%; Score 36.8; DB 12; 45.3%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                      09-AUG-2001.
(HUVA-) HUMAN GENOME SCI INC.
(HIVA-) HUMAN GENOME SCI INC.
(ery Match 52.8; Score 36.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADA98754 standard; DNA; 54877 BP.
Human secreted protein-related DNA sequence #347.
WO2003004623-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MCACCOLORY
16-7AA-2003.
(HUAA-) HUMAN GENOME SCI INC.
6-ry Match
5.2%; Score 36.8; DB 8;
--- Cimilarity 50.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABX08336 standard; DNA; 1691080 BP.

3 Human phosphodiesterase 4D (PDB4D) gene.

Query Match
5.2%; Score 36.8; DB 6;
Best Local Similarity 45.3%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADJ25985 standard; DNA; 1691139 BP.
Human phosphodiesterase 4D (PDE4D) genomic DNA.
US2004014099-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADN97989 standard; DNA; 1691138 BP.
Human phosphodiesterase 4D genomic sequence.
WO2004028341-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADOS0281 standard; DNA; 1691134 BP.
Human phosphodiesterase 4D (PDE4D) gene.
US2004091865-Al.
13-MAY-2004.
AAK86025 standard; DNA; 54863 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity 50.6%; Prec
RESULT 1273
ID ABZ67791 standard; DNA; 54877 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     03-OCT-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     08-APR-2004.
(DECO-) DECODE GENETICS EHF.
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                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1271
D ABZ74225 standard; DN DE Secreted protein gene
PN W0200277013-A2.
PA (HUMA-) HUMAN GENOME
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RESULT 1272
                                                                                                    Query Match
Best Local Similarity
RESULT 1270
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Best Local Similarity
RESULT 1274
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                                                                                                                                                                                                                                                                      Query Match
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Best Local S
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03-JAN-2002
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WO200168911-A2.
                                                                                                                                                                                                                          ABLO1896 standard; cDNA; 4755 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 170.
WO20011042-A2.
Z7-SED-2001.
(PEKE ) PE CORP NY.
                                              5.1%; Score 36.6; DB 12; Length 4018; 57.4%; Pred. No. 1.3e+02;
                                                                                                                                                                            5.1%; Score 36.6; DB 10; Length 4316; 52.3%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 7000;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 7657;
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                                                                                                                                                                                                                                                                                                        Length 4755
                                                                                                                                                                                                                                                                                                                                                   ADF74255 standard; DNA; 7000 BP.
Genomic DNA sequence of the pAA plasmid of EAEC strain 042.
US2003180315-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human chemically pretreated gene sequence #71 strand WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAS63351 standard; DNA; 7025 BP. Chemically pretreated metabolism associated gene #46 WO200176451-A2.
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Human immune system associated gene SEQ ID NO: 1870.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABLA9022 standard; DNA; 7657 BP.
Human immune system associated gene SEQ ID NO: 1995.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABL32983 standard; DNA; 7143 BP.
Human immune system associated gene SEQ ID NO: 956.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                 5.1%; Score 36.6; DB 10; 49.2%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          5.1%; Score 36.6; DB 6; 54.0%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 36.6; DB 6;
Pred. No. 1.4e+02;
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(EPIG-) EPIGENOMICS AG.

5.1%; Score 36.6; DB 6;

ery Match 5.1%; Pred. No. 1.4e+02;
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                                                                                                                                                                                                                                                                                                        Score 36.6; DB 4;
Pred. No. 1.3e+02;
                                                                                                               Pretreated genomic DNA region 148.
WO2003072821-A2.
04-SEP-2003.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAS45477 standard; DNA; 7657 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AEK40060 standard; DNA; 7025 BP
                                                                                                 DNA; 4316 BP.
                   01-JAN-2004.
(UYOR-) UNIV OREGON HEALTH SCI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.1%;
                                                                                                                                                                                                                                                                                                          5.18;
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(EPIG-) EPIGENOMICS AG.
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(BPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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RESULT 1295
ID ABL34022 standard; DN
DE Human immune system 6
PN WO2002028-A2.
PD 03-JAM-2002.
PA (EPIG-) EPIGENOMICS A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 1296
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1293
                                                                                                                                                                                         Best Local Similarity RESULT 1289
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                                                   Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
                                                                                                 ADB54224 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                       (NATA/) NATARO J.
    US2004003422-A1.
                                                                                                                                                                                                                                                                                                                                                                                                        25-SEP-2003
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                                                                                                                                                                                                                                                                                                                                      RESULT 1290
ID ADF7425E
DE Genomic
PN US200318
PD 25-SEP-7
PA (NATA/)
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AAD03729 standard, DNA, 10640 BP.
AAD03729 standard, DNA 10640 BP.
P. falciparum telomerase reverse transcriptase (TERT) full length gene.
WO200127287-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        האסשיטון standard; DNA; 11260 BP.
Chemically pretreated genomic DNA associated with cell cycle #10.
WO200168911-A2.
20-SEP-2001.
                                                                                                                                                                                                                         AAS45452 standard; DNA; 9515 BP.
Chemically pretreated genomic DNA associated with cell cycle #79-
W0200168911-A2.
20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
EY MAtch
st Local Similarity 52.3%; Pred. No. 1.5e+02;
                                                                                           NO 19142
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     261
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                   Length 8143;
                                                                                                                                                                  Length 8242,
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Human polynucleotide associated with DNA replication SEO
W0200177377-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABNB0244 standard; DNA; 9515 BP.
Human chemically modified disease associated gene SBQ ID
WO200200927-A2.
                                                                          ABLO8220 standard; cDNA, 8242 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID
W0200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    725
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(BPIG-) EPIGENOMICS AG.
(EPIG-) Aduch 5.1%; Score 36.6; DB 6;
(ery Match 52.3%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 36.6; DB 6;
Pred. No. 1.6e+02;
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Pred. No. 1.6e+02;
                                                                                                                                                                  Score 36.6; DB 4;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 36.6; DB 6;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 36.6; DB 6;
Pred. No. 1.6e+02;
PA (EPIG-) EPIGENOMICS AG.
Ouery Match
Best Local Similarity 47.6%; Pred. No. 1.5e+02;
RESULT 1297
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 11131 BP.
DNA repair gene fragment#18
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                        DNA transcription associated genomic DNA #91.
WO200192565-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity 46.3%; Pred
RESULT 1302
ID ABL32752 standard; DNA; 11131 BP.
                                                                                                                                                                                                                                                                                                                                                                           BP.
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03-JAN-2002
(EPIG-) EPIGENOMICS AG.
EPIG-) EPIGENOMICS AG.
5.1%;
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Query Match 5.1%;
Best Local Similarity 58.9%;
RESULT 1304
                                                                                                                                                                      5.18;
                                                                                                                                                                                                                                                                                                                                                                         ABK28307 standard; DNA; 9515
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(RERE-) RES & DEV INST INC.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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RESULT 1303
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL9226 standard;
Chemically treated
WO200181622-A2.
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RESULT 1301
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RESULT 1299
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RESULT 1298
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us-09-989-293a-376.rng.spdi

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S. epidermidis open reading frame nucleotide sequence SEQ ID NO:1949.
WO200134809-A2.
                                                                                                                                                                                        AAS46789 standard; DNA; 61020 BP.
Tumour suppressor gene derived chemically modified sequence #514 W0200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%; Score 36.6; DB 12; Length 191395; 68.0%; Pred. No. 2.9e+02;
     5.1%; Score 36.6; DB 10; Length 52242; 52.3%; Pred. No. 2.2e+02;
                                                                                                                                                   Length 52242;
                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 113515;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 159095;
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                                                                                                                                                                                                                                                                                             Length 61020;
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                                                                                                                                                                                                                                                                                                                   Human immune system associated gene SEQ ID NO: 2148. WO200200928-A2.
                                                                                                                                               Score 36.6; DB 10;
Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 36.6; DB 6;
Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 5 1%: Score 36.4; DB 5;
                                                                                                                                                                                                                                                                                           Score 36.6; DB 4;
Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.1%; Score 36.6; DB 8; 68.0%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 36.6; DB 4;
Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.1%; Score 36.4; I
45.6%; Pred. No. 83;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.1%; Score 36.4; I
45.6%; Pred. No. 83;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADL08126 standard; DNA; 191395 BP.
Human gene associated with low HDL-C PAI2.
US2004043389-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
5.1%; Score 36.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human ovarian cancer DNA; 318 BP.

WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human ovarian cancer DNA marker #5902.
WO200170979-A2.
                                                      ADE95914 standard; DNA; 52242 BP.
Human MDM2 gene genomic DNA sequence.
WO2003039484-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABZ80818 standard; DNA; 159095 BP.
Human PAI-2 polymorphism PAI2u1.
WO2003007801-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAH88704 standard; DNA; 160755 BP.
Human DNA sequence SEQ ID 544.
WO200151659-A2.
                                                                                               Wozucz--
15-MAY-2003.
(SAGR-) SAGRES DISCOVERY.
Gerv Match ---11arity 52.3%;
                                                                                                                                                                                                                                            WOZYCZ
20-SED 2001.
(EPIG-) EPIGENOMICS AG.
5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                     03-JAN-ZUUG.
(EPIG-) EPIGENOMICS AG.
SIY, MATCh 5.1%;
SIY ---? Cimilarity 45.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADI73160 standard; DNA; 318
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAH53278 standard; DNA; 342
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 04-MAR-2004.
(VITI-) VITIVITY INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           30-JAN-2003.
(VITI-) VITIVITY INC.
                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 1317
   Query Match
Best Local Similarity
RESULT 1315
                                                                                                                                               Query Match
Best Local Similarity
RESULT 1316
                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 1318
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 1323
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         19-JUL-2001.
(GEST ) GENSET.
                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABK31511 standard; DNA; 47108 BP.
Signal transduction associated gene modified complementary DNA #177.
WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 авы19792 standard; DNA; 33472 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 10849.
                  Score 36.6; DB 4; Length 11260; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 11996;
                                                                                                                                                              Length 11260;
                                                                                                                                                                                                                                                                                                          Length 11260;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AASO2354 standard; DNA; 27048 BP.
Porcine Gal alpha(1,3) galactosyl transferase gene introns 3-0
WO200123541-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 27048
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 33472
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 47108
                                                                                                                                                                                                                                 Human chemically modified disease associated gene SEQ ID NO WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADA02666 standard; DNA; 52242 BP.
Human MDM2 carcinoma associated gene, SEQ ID NO:1184.
820303057146-A2.
17-JUL-2003.
                                                                                                                                    (EPIG-) EPIGENOMICS AG. 5.1%; Score 36.6; DB 6; ry Match t. Local Similarity 44.8%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                   WOZUWI...
18-OCT-2001.
(EPIG- PEIGENOMICS AG.
Ery Match
Ery Match
5.1%; Score 36.6; DB 6;
Fry Match
7. "milarity 48.0%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               58.9%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.1%; Score 36.6; DB 4; 56.1%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.1%; Score 36.6; DB 9; 52.3%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                        Score 36.6; DB 6;
Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.1%; Score 36.6; DB 6; 48.4%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 36.6; DB 2;
Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                      ABL34493 standard; DNA; 11996 BP.
Human metastasis associated gene SEQ ID NO: 46.
WO200177376-A2.
                                                                    ABK28153 standard; DNA; 11260 BP.
DNA transcription associated genomic DNA #14
06-DEC-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Tomato pest resistance Mi gene (copy 1) NO9915171-A1.
                                                                                                                                                                                                                  ABN80038 standard; DNA; 11260 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA; 52242 BP
                  5.1%;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.14;
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(ALEX-) ALEXION PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (SAGR-) SAGRES DISCOVERY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                30-JAN-2003,
(SAGR-) SAGRES DISCOVERY
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (REGC ) UNIV CALIFORNIA
(EPIG-) EPIGENOMICS AG.
                             Best Local Similarity RESULT 1306
                                                                                                                                                                             Best Local Similarity RESULT 1307
                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 1308
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1309
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Best Local Similarity
RESULT 1312
ID AAV26084 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 1310
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(PEKE ) PE CORP NY.
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Best Local Similarity
RESULT 1311
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1313
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RESULT 1314
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human MDM2 gene.
WO2003008583-A2.
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                  Query Match
                                                                                                                                                            Query Match
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(GETH ) GENENTECH INC.
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RESULT 1337
                                                                                                                                                                                            ABN92001 standard; DNA; 408 BP.
Staphylococcus epidermidis ORF nucleic acid sequence SEQ ID NO:1464.
Us6380370-B1.
                  LT 1324
ABX47837 standard; cDNA; 344 BP.
Bovine EST associated with lactation/muscle/fat deposition #13002.
US2002137139-A1.
26-SEP-2002.
(WATH) MATHIALAGAN N.
(TAON) TAO N.
(WARR) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.1%; Score 36.4; DB 10; Length 1230; 61.7%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 1256;
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                                                                                                                                                                                                                                                                      SUCRION-) GENOME THERAPEUTICS CORP.

ry Match
Local Similarity 56.8%; Pred. No. 88;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 6; Length 462;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          5.1%; Score 36.4; DB 6; Length 926; 53.5%; Pred. No. 1e+02;
                                                                                                                                                                DB 8; Length 344;
                                                                                                                                                                                                                                                                                                                                                                                                                       Length 435;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              570.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABT40613 standard; DNA; 479 BP.
Toxicity modelling related rat gene SEQ ID No 315.
WC200295000-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL35041 standard; cDNA; 926 BP. Murine cDNA isolated from skin cells SEQ ID NO: WO200190357-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.1%; Score 36.4; DB 3; 58.2%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                          4
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Human protein kinase encoding cDNA SEQ ID NO:37.

W0200300901-A2.

(95-5AN-2003.

(DECO-) DECODE GENETICS EHF.
                                                                                                                                                                                                                                                                                                                                                                                                                       DB
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47.7%; Pred. No. 91;
                                                                                                                                                              5.1%; Score 36.4; 3 52.7%; Pred. No. 85;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 y Match 5.1%; Score 36.4; Local Similarity 44.2%; Pred. No. 90;
                                                                                                                                                                                                                                                                                                                                                                                                                       5.1%; Score 36.4;
51.2%; Pred. No. 89;
   56.8%; Pred. No. 85;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABN93640 standard; DNA; 462 BP.
Gene #138 used to diagnose liver cancer.
WC200229103-A2.
                                                                                                                                                                                                                                                                                                                                     AA191807 standard; cDNA; 435 BP.
Human polynucleotide SEQ ID NO 11867.
WO200164835-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABN81046 standard; DNA; 1256 BP. Shrimp polynucleotide SEQ ID NO 50. WO200034476-A2. I5-JUN-2000 (TUFT) TUFTS COLLEGE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            29-NOV-2001.
(GENE-) GENESIS RES & DEV CORP LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADO20394 standard; DNA; 1506 BP.
Human PRO polynucleotide #760.
WO2004043361-A2.
27-MAY-2004.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  28-NOV-2002.
(GENE-) GENE LOGIC INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    11-APR-2002.
(GENE-) GENE LOGIC INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2002.
2002.
2002.
4002.
Ary Match
Best Local Similarity 47
RESULT 1329
ID ABL35041 standar
DE Murine CDNA
PD 29-20
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Best Local Similarity
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                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1325
                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
     Best Local Similarity
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                               30-APR-2002
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ID ABT40613
DE Toxicity
PN WC20029
PD 28-NOV-2
PA (GENE-)
                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 1327
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                       RESULT
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ADM91294 standard; DNA; 1902 BP.
DNA homologue of a human DNA sequence encoding a protein SeqID 628.
WO2004020595-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 2308;
                                                                                                                                                                                                                                                                                                                                                                                          yuery match 5.1%; Score 36.4; DB 12; Length 1902;
Best Local Similarity 61.7%; Pred. No. 1.26+02;
RESULT 1335
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 2308;
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A (HELL -) HELIX RES INST.

Query Match
5.1%; Score 36.4; DB 4; Length 2308;

Best Local Similarity 51.2%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH54088 standard; DNA; 3005 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:3452.
WO200134809-A2.
  Length 1506;
                                                                                                                                                                                   Length 1559;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             T7-MAY-2001.
(GLAX ) GLAXO GROUP LTD.

ery Match

5.1%; Score 36.4; DB 4; Length 3005;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 2000;
Query Match
Best Local Similarity 56.8%; Pred. No. 1.2e+02;
RESULT 1333 standard; DNA; 1559 BP.
DE Human soft tissue sarcoma-upregulated DNA - SEQ ID 8273.
PD 10-UIN-2004.
PA (PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACC72733 standard; cDNA; 2308 BP.
Human cancer related protein encoding cDNA SEQ ID NO:72.
WO2003025138-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match 5.1%; Score 36.4; DB 12; Beet Local Similarity 51.2%; Pred. No. 1.3e+02; RESULT 1339
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.1%; Score 36.4; DB 10; 51.2%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (FIVE-) FIVE PRIME THERAPEUTICS INC.

(FIVE-) FIVE PRIME THERAPEUTICS INC.

5.1%; Score 36.4; DB 12;

FINE Score 36.4; DB 12;

FINE Score 36.4; DB 12;

FINE Score 36.4; DB 12;
                                                                                                                                                                       Luery match 5.1%; Score 36.4; DB 12; Best Local Similarity 56.8%; Pred. No. 1.2e+02; RESULT 1334
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.1%; Score 36.4; DB 8; 55.6%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADL91782 standard; cDNA; 2308 BP.
Human PRO69889 encoding cDNA SEQ ID NO:3.
                                                                                                                                                                                                                                                                                                                           11-MAR-2004.
(FIVE-) FIVE PRIME THERAPEUTICS INC.
(RIKE-) RIKEN INST PHYSICAL & CHEM RES.
(DNAF-) DNAFORM KK.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAH16252 standard; cDNA; 2308 BP.
Human cDNA sequence SEQ ID NO:15093.
EP1074617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADA72008 standard; DNA; 2000 BP.
Rice gene, SEQ ID 5333.
WO20030008998-Al.
(33-JAN-2003.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAX25320 standard; cDNA; 3331 BP. Mouse TNF receptor TRL cDNA. W09915663-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AD035529 standard; DNA; 2506 BP.
Novel mouse gene sequence #202.
WO2004046310-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 27-MAR-2003.
(EOSB-) EOS BIOTECHNOLOGY INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1341
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
RESULT 1336
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 1338
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25-MAR-2004.
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AAS45321 standard; DNA; 9293 BP.
Chemically pretreated complementary DNA associated with cell cycle #13.
WO200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 10907;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.1%; Score 36.4; DB 12; Length 10907; 61.7%; Pred. No. 1.7e+02;
                            Length 8979;
                                                                                                                                                                                     Length 8979;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 8979
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 9293;
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PA (EPIG-) EPIGENOMICS AG.

Query Match

S.1%; Score 36.4; DB 6; Length 897

Best Local Similarity 51.9%; Pred. No. 1.7e+02;

RESULT 1333

ID AAS61178 standard; DNA; 8979 BP.

DE Human gene regulation-associated gene oligonucleotide #133.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 1356

ID ABK28160 standard, DNA, 9293 BP.

DE DNA transcription associated complementary genomic DNA #17

PN W0200192565-A2.
                                                                                                        #57
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABK39974 standard; DNA; 9293 BP.
Human chemically pretreated gene sequence #28 strand
WO200202806-A2.
                                                                                                                                                                                                                                           DNA; 8979 BP.
cell signalling DNA sequence#61
                                                                                                      Signal transduction associated gene modified DNA WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 36.4; DB 12;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         PD 18-OCT-2001.

PA (EPIG-) EPIGENOMICS AG.

Querry Match 5.1%; Score 36.4; DB 6;
Best Local Similarity 51.9%; Pred. No. 1.7e+02;

RESULT 1354
                      Score 36.4; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                 Score 36.4; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADJ62879 standard; DNA; 10907 BP.
Human p21 WAF-1 gene related to cancer treatment.
US2003228285-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 36.4; DB 4;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 36.4; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 36.4; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAH48024 standard; DNA; 14041 BP.
Internal control B19c #1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA; 10907 BP.
                                                                                     DNA; 8979 BP.
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                    Query Match
Best Local Similarity 51.9%;
RESULT 1351
                                                                                                                                        PD 03-JAN-2002.
PA (EPIG-) EPIGENOMICS AG.
Query Match 5.1%;
Best Local Similarity 51.9%;
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Local Similarity 54.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ACF57521 standard; DNA; 1090
Human p21 DNA SEQ ID NO: 22.
WQ2003093303-A1.
13-NOV-2003
(TEXA ) UNIV TEXAS SYSTEM.
    (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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                                                                                                                                                                                                                                    ABL70231 standard;
Chemically treated
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 1358
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RESULT 1359
                                                                               ABK31270 standard;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (HUNG/) HUNG M.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   WO200146463-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        11-DEC-2003
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                                                                                                                                                                                                  Length 3655;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAD46980 standard; DNA; 5917 BP.
Plasmodium falciparum merozoite surface protein-1 (MSP-1) DNA
WO200270542-A2.
                                          Length 3331;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 5917;
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                                                                                                                                                                                                                                                                                                                                                                                      RESULT 1344
ID AANSOS30 standard; DNA; 5760 BP.
DE Sequence encoding the P195 protein of Plasmodium falciparum.
PN EP154454-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABK28370 standard; DNA; 6167 BP.
DNA transcription associated complementary genomic DNA #122.
WO20192565-A2.
(EPIG-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAS64432 standard; DNA; 5986 BP.
Human gene regulation-associated gene oligonucleotide #387.
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DNA; 6935 BP.
DNA repair gene fragment complementary
                                                                                                                    Human soft tissue sarcoma-upregulated DNA - SEQ ID 7424
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Signal transduction associated gene modified DNA #171 WO200200926-A2.
                                                                                                                                                                                                                                                   ABL32957 standard; DNA; 5182 BP.
Human immune system associated gene SEQ ID NO: 930.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL32784 standard; DNA; 8979 BP.
Human immune system associated gene SEQ ID NO: 757.
WO200200928-A2.
                                                                                                                                                                                              5.1%; Score 36.4; DB 12; 51.2%; Pred. No. 1.4e+02;
                                      5.1%; Score 36.4; DB 2; 59.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                      03-3-002.
03-3-002.
(EPIG-) EPIGENOMICS AG.
(ery Match
(ery Match
56.8%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        5.1%; Score 36.4; DB 6; 45.9%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 36.4; DB 6;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.1%; Score 36.4; DB 6; 50.6%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.1%; Score 36.4; DB 1; 45.9%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        5.1%; Score 36.4; DB 6; 49.5%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (SELI-) ST ELIZABETH'S MEDICAL CENT INC.
                (MILL-) MILLENNIUM BIOTHERAPEUTICS INC
                                                                                                                                                                           (PROT-) PROTEIN DESIGN LABS INC.
                                                                                                ADQ24604 standard; DNA; 3655 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABK31498 standard; DNA; 5986 BP.
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
MATCH ''narity 49.5%;
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(WELL ) WELLCOME FOUND LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                01-NOV-2001.
(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                     Best Local Similarity RESULT 1342
                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 1343
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Best Local Similarity
RESULT 1345
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Best Local Similarity
RESULT 1346
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Best Local Similarity
RESULT 1347
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RESULT 1349
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Chemically treated
WO200181622-A2.
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01-APR-1999
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                                      Query Match
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(EPIG-) EPIGENOMICS AG.
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               Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL70374 standard; DNA; 15951 BP. Chemically treated cell signalling DNA sequence complementary to#132.WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL70514 standard; DNA; 15674 BP. Chemically treated cell signalling DNA sequence complementary to#202. WC200202807-A2.
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                          Length 14041;
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                                                                                                                                    Length 14551;
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Human immune system associated gene SEQ ID NO: 1654.
00210200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL32654 standard; DNA; 17211 BP.
Human immune system associated gene SEQ ID NO: 627.
WO200200928-A2.
03-JAN-2002.
                                                                                                                                                                         ABL62905 standard; DNA; 15000 BP.
Breast cancer related gene sequence SEQ ID NO:1242.
WO200194629-A2.
                                                                                                                                                                                                                                                                            ABL32363 standard; DNA; 15674 BP.
Human immune system associated gene SEQ ID NO: 336
WO200200928-A2.
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10-JAN-2002.

(EPIG) EPIGENOMICS AG.

5.1%; Score 36.4; DB 6;

ery Match

5.1%; Pred. No. 1.9e+02;
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                            5.1%; Score 36.4; DB 4; 16.0%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                           OB3-UAN-ZUUS.
(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.4; DB 6;
ry Annal Similarity 47.4%; Pred. No. 1.9e+02;
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Pred. No. 1.9e+02;
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Pred. No. 1.9e+02;
                                                                                                                                  Score 36.4; DB 6;
Pred. No. 1.9e+02;
                                                                                                                                                                                                                                          5.1%; Score 36.4; DB 6; 47.1%; Pred. No. 1.9e+02;
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Human metastasis associated gene SEQ ID NO: 134
WO200177376-A2.
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Pred. No. 1.9e+02;
                                                                  ABL34585 standard; DNA; 14551 BP.
Human metastasis associated gene SEQ ID NO: 138
WO200177376-A2.
                                                                                                                                                                                                                                                                                                                                                                                    ABL34477 standard; DNA; 15674 BP.
Human metastasis associated gene SEQ ID NO: 30.
WC200177376-A2.
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%;
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(EPIG-) EPIGENOMICS AG.
127 Match 5.1%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                                                          18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
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(AVAL-) AVALON PHARM.

Query Match

Best Local Similarity
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RESULT 1365
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Best Local Similarity
RESULT 1366
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RESULT 1361
                                       Best Local Similarity
RESULT 1360
    28-JUN-2001.
(BAXT ) BAXTER AG.
Query Match
                                                                                                                                                                                                                                                                                                                          03-JAN-2002
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ID NO:25435
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Invertebrate foraging behaviour associated human DNA sequence #12.
01-AUG-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAS46745 standard; DNA; 38342 BP.
Tumour suppressor gene derived chemically modified sequence #469
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 29993;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 38342;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human testicular antigen encoding DNA fragment SEQ ID NO: 2665 WO200155317-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 5.1%; Score 36.4; DB 4; Length 29163;
                                                                                                                                                          Length 19734;
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Length 17211;
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                                                                                                                                                                                                                                                                                                                                                                             AAL05121 standard; DNA; 29163 BP.
Human reproductive system related antigen DNA SEQ ID NO: 7809.
W0200155320-A2.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 29163;
                                                                                                                                                                                                                                     Human immune/haematopoietic antigen genomic sequence SEQ
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Signal transduction associated gene modified DNA #175 WO200200926-A2.
                                                             ABL33933 standard; DNA; 19734 BP.
Human immune system associated gene SEQ ID NO: 1906
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADB37660 standard, DNA, 29993 BP.
Human chemically pretreated EYA4 gene SEQ ID NO:2.
WO2003072812-A2.
                                                                          Human immuse ...
W0200200928-A2.
W0200200928-A2.
(G) JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.4; DB 6; ...
Match ...rity 45.7%; Pred. No. 2e+02;
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Pred. No. 2.2e+02;
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PD 20-SEP-2001.

PA (EPIG-) EPIGENOMICS AG.

Query March

Best Local Similarity 49.0%; Pred. No. 2.3e+02;
Score 36.4; DB 6;
Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                 (HUMA-) HUMAN GENOME SCI INC.

Y Match
5.1%; Score 36.4; DB 4;

L Local Similarity 61.7%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 5.1%; Score 36.4; DB 4;
Best Local Similarity 61.7%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 36.4; DB 6;
Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 36.4; DB 8;
Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human ribosomal DNA complete repeating unit. US2002160970-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABK31506 standard; DNA; 38342 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADA14747 standard; DNA; 42998 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABL98013 standard; DNA; 29163 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     04-SEF-2005.
(EPIG-) EPIGENOMICS AG.
FY MATCh 5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.1%;
    5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        31-OCT-2002.
(HADL/) HADLACZKY G.
(SZAL/) SZALAY A A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity
RESULT 1376
                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 1371
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best_Local Similarity
RESULT 1374
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
                                                                                                                                                                             Best Local Similarity
RESULT 1370
                  Best Local Similarity RESULT 1369
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 04-SEP-2003
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DE NO E

us-09-989-293a-376.rng.spdi

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CN1364918-A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10-JUL-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
       RESULT 1387
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL03438 standard; cDNA; 76574 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 4796
WO200171042-A2.
                                                                                                                                                                               Length 42999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 54786;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 54786;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 96587;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 95400,
                                                                                                                                                                                                                                                                                                                       Length 54786;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%; Score 36.4; DB 9; Length 96589; 45.7%; Pred. No. 2.8e+02;
                     Length 42999;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 76574;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 91000;
                                                                                                                                                                                                                              ADA02705 standard; DNA; 54786 BP.
Mouse Zfhx1b carcinoma associated gene, SEQ ID NO:1223
WO2003057146-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human JAK2 carcinoma associated gene, SEQ ID NO:1430 WO2003057146-A2.
                                                                                                                                                                             5.1%; Score 36.4; DB 10;
61.7%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADPO8388 standard; DNA; 95400 BP.

Human laminin alpha 4 (LAMA4) genomic DNA.

ery Match

5.1%; Score 36.4; DB 12;

st Local Similarity 45.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 36.4; DB 10;
Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.1%; Score 36.4; DB 10; 61.7%; Pred. No. 2.5e+02;
(NEUR-) NEUROSCIENCES RES FOUND INC.
5.1%; Score 36.4; DB 6;
it Local Similarity 61.7%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.1%; Score 36.4; DB 4;
53.5%; Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Alici326 standard; DNA, 91000 BP.

Human farnesoid X receptor (FXR) DNA #2.

W02003044167-A2.
30-MAY-2003.

(ISIS PHARM INC.
5.1%; Score 36.4; DB 9;

St Local Similarity 61.7%; Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                  5.1%; Score 36.4; DB 9; 61.7%; Pred. No. 2.5e+02;
                                                                      AAD61411 standard; DNA; 42999 BP.
Human ribosomal DNA (rDNA) repeat region.
US2003101480-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Mouse Zhxlb gene genomic DNA sequence. WO2003039484-A2.
                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity 61.7%; Pred
RESULT 1380
ID ADB72443 standard; DNA; 54786 BP.
DE Mouse Zihxlb gene.
PN WO2003008583-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADC85392 standard; DNA; 96587 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADA02912 standard; DNA; 96589 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human Jak2 genomic sequence.
WO2003045230-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        05-JUN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   17-JUL-2003, (SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                     30-JAN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                   (SAGR-) SAGRES DISCOVERY
                                                                                                                     29-MAY-2003.
(HADL/) HADLACZKY G.
(SZAL/) SZALAY A A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   27-SEP-2001.
(PEKE) PE CORP NY.
                   Query Match
Best Local Similarity
RESULT 1378
                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1379
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1382
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1383
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RESULT 1384
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Best Local Similarity
RESULT 1385
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                     17-JUL-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
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ADF13118 standard; DNA; 161652 BP. Hypermethylation site in human breast cancer CpG island locus HBC-48. US2003129602-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABX48177 standard; cDNA; 286 BP.
Bovine EST associated with lactation/muscle/fat deposition #13342.
US2002137139-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 161652;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.1%; Score 36.4; DB 12; Length 161652; 47.4%; Pred. No. 3.1e+02;
                                                                                                             Length 96589;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 106416;
                                                                                                                                                                                                                                                                                                        Length 96589
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 110000;
                                                                                                                                                                                                                                                                                                                                                              ABLI8718 standard, DNA, 106416 BP.
Drosophila melanogaster genomic polynuclectide SEQ ID NO 7627, WWO200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 251,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 286;
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ID AAL14832 standard; CDNA; 287 BP.

E Human breast cancer expressed polynucleotide 7289.

PN W02019128-A2.

PD 19-JUL-2001.
                                                                                              Query Match 5.1%; Score 36.4; DB 10;
Best Local Similarity 45.7%; Pred. No. 2.8e+02;
RESULT 1388
                                                                                                                                                                                                                                                                                                     Score 36.4; DB 12;
Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 36.4; DB 12;
Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB 10;
                                                                                                                                                                                            Human carcinoma associated (CA) nucleic acid #88 US2004072154-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.1%; Score 36.4; DB 4; 47.7%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%; Score 36.4; DB 6; 53.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity 47.7%; Pred. No. 2.8e+02
RESULT 1390
ID ABA90521 standard; DNA; 2365589 BP.
DE Genomic sequence of Lactococcus lactis IL1403.
PN FR2807446-A1.
PD 12-OCT-2001.
PA (INRG) INRA INST NAT DECH ADMINISTRA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match 5.1%; Score 36.2; I
Best Local Similarity 51.6%; Pred. No. 88;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.1%; Score 36.2; 52.3%; Pred. No. 91;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABX94407 standard; cDNA; 251 BP.
Rice endosperm-associated EST cDNA SEQ ID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AD137264 standard; DNA; 161652 BP.
Hypermethylation in cancer (HBC) locus-48.
US6605432-Bl.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (INRG ) INRA INST NAT RECH AGRONOMIQUE.
ADB72650 standard; DNA; 96589 BP. Human JAK2 gene. Mo200300808583-A2. 30-JAN-2003. (SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                          DNA; 96589 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.1%;
                                                                                                                                                                                                                                                                                                     5.1%;
                                                                                                                                                                                                                                                         (MORR/) MORRIS D W. (ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              21-AUG-2002.
(UYZH-) UNIV ZHEJIANG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (BYAT/) BYATT J C. (MATH/) MATHIALAGAN N.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          12-AUG-2003.
(UMOR ) UNIV MISSOURI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 1391
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 1392
                                                                                                                                                                        ADM74507 standard;
                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 1389
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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5; Length 473;

Length 538;

Length 538;

Length 725;

Length 496;

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Arabidopsis thaliana stress regulated gene SEQ ID NO 5202 WO200216655-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAX37532 standard; cDNA; 725 BP.
Human secreted protein cDNA fragment containing gene 82
WO9918208-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 1409
ID ADC86904 standard; DNA; 919 BP.
DE Human GPCR gene SEQ ID NO:1357.
PN BP1270724-A2.
PD C2-JAW-2003.
PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
Query Match
5.1%; Score 36.2; DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Nanoarchaeum equitans cancer-associated (CA) gene WO2003093434-A2.
                                                                          AAL16526 standard; cDNA; 496 BP,
Human breast cancer expressed polynucleotide 8983.
WO200151628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAD37062 standard; cDNA; 2396 BP.
Influenza B virus/vienna/1/99/ca PB2 mutant cDNA.
WO200224876-A2.
28-MAR-2002.
                                                                                                                                 19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
SATA STATE SIMILARITY 49.4%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                    PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Query Match 5.1%; Score 36.2; DB 5;
Best Local Similarity 51.4%; Pred. No. 1e+02;
RESULT 1407
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   D 15-APR-1999.

4 (HUMA-) HUMAN GENOME SCI INC.

Query Match

5.1%; Score 36.2; DB 2;

Best Local Similarity 52.3%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      28-FEB-2002.
(SCRI ) SCRIPPS RES INST.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
(SYGN ) SYNGENTA FARTICIPATIONS AG.
17 Match
11 Local Similarity 53.6%; Pred. No. 1.40+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.1%; Score 36.2; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5.1%; Score 36.2; DB 8; 45.8%; Pred. No. 1.3e+02;
                Score 36.2; DB
Pred. No. 1e+02;
                                                                                                                                                                                   Best Local Similarity 49.4%; Pred. No. RESULT 1406

ID ADL37492 standard; DNA; 538 BP.

DE Human ovarian cancer DNA marker #11382.
                                                                                                                                                                                                                                                                                                                                                                                         AD172348 standard; DNA; 538 BP.
Human ovarian cancer DNA marker #5090.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Prokaryotic essential gene #4779 W0200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADK16276 standard; DNA; 1062 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ACA23122 standard; DNA; 1587 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABZ17397 standard; DNA; 2000 BP.
                  5.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                J. NAT INST , J. OSC., CENT ADVANC , LOCAL Similarity , RESULT 1410 ID ADK16776 standar DE Nanoarchaen PN WO20030^C PD 13-**
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       13-NOV-2003.
(DIVE-) DIVERSA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
RESULT 1412
                  Query Match
Best Local Similarity
RESULT 1405
                                                                                                                                                                                                                                                                                                27-SEP-200
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Si
RESULT 1408
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          LT 1399
AALI189 standard; CDNA; 393 BP.
Human breast cancer expressed polynucleotide 3646.
Human breast cancer expressed polynucleotide 3646.
HUMC200151628-A2.
19-UUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.1%; Score 36.2; DB 4; Length 393;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.1%; Score 36.2; DB 5; Length 469;
ery Match
5.1%; Pred. No. 1e+02;
                                                                                                                                                      23.AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
5.1%; Score 36.2; DB 5; Length 325;
                                                                                                                                                                                                                                                                                                                                     (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Query Match 5.1%; Score 36.2; DB 4; Length 367;
Best Local Similarity 49.4%; Pred. No. 96;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 369;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 4; Length 394;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 461;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 464;
                                     Length 287;
                                                                                                                                                                                                                                                       AAL25369 standard; cDNA; 367 BP.
Human breast cancer expressed polynucleotide 17826.
WP0200151628-A2.
19-JUL-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           14-JUN-2001.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Query Match 5.1%; Score 36.2; DB 4;

Best Local Similarity 43.9%; Pred. No. 96;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
5.1%; Score 36.2; DB 5;
r Innal Similarity 49.6%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.1%; Score 36.2; DB 5; 50.9%; Pred. No. 1e+02;
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 5 1%. Score 36.2; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                               AAH69544 standard; cDNA; 369 BP.
Human cervical cancer marker nucleic acid 818.
WC200142467-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABV55326 standard; cDNA; 464 BP.
Human prostate expression marker cDNA 55317.
WO200160860-A2.
                                                                                            ABV59017 standard; cDNA; 325 BP.
Human prostate expression marker cDNA 59008.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human prostate expression marker cDNA 55188.
WO200160860-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABV09921 standard; cDNA; 473 BP.
Human prostate expression marker cDNA 9912.
WC200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.1%; Score 36.2; D 51.0%; Pred. No. 97;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        5.1%; Score 36.2; I
49.2%; Pred. No. 97;
                                         5.1%; Score 36.2;
40.9%; Pred. No. 91;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AA187163 standard; cDNA; 394 BP.
Human polymucleotide SEQ ID NO 7223.
WO200164835-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABV55197 standard; cDNA; 469 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABV08266 standard; cDNA; 461 BP.
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2001.

LLL-) MILLENNIUM

LY MAtch

Best Local Similarity

RESULT 1404

ID ABV0921 stand*

DE Human prost

PN WC20016*

PD 23-*

PA
                                                                                                                                                                                                                Best Local Similarity RESULT 1397
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Best Local Similarity
RESULT 1402
                                         Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
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Best Local Similarity
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(HYSE-) HYSEQ INC.
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                                                                                                                                                                                                 Query Match
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ID AAH69544
DE Human of
PN WC200142
PD 14-JUN-7
PA (MILL-)
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ID ABV0826

DE Human P.

PN W020016

PD 23-AUG--

PA (MILL-)
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Length 1062;

Length 1587;

Length 2000;

Length 919;

Query Match

Query Match

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14.
                                           ABL49314 standard; DNA; 6113 BP.
Human polynucleotide associated with DNA replication SEQ ID NO-WO200177377-A2.
         Length 6113;
                                                                                                                                                                                                                                                                                         Length 6121;
                                                                                                                                                Length 6113;
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Human gene regulation-associated gene oligonucleotide
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                            #22
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABK40057 standard; DNA; 7441 BP.
Human chemically pretreated gene sequence #70 strand
WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 #39
                                                                                                                                                                                               ABL33974 standard; DNA; 6121 BP.
Human immune system associated gene SEQ ID NO: 1947.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL/U167 standard; DNA; 6154 BP.
Chemically treated cell signalling DNA sequence#29.
WQ200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              457
                                                                                                                                                                                                                                                                                                              Signal transduction associated gene modified DNA (03-UAN-2002.)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Chemically pretreated metabolism associated gene WO200176451-A2.
         Score 36.2; DB 6;
Pred. No. 1.7e+02;
                                                                                                                                                                                                                                             03-JAN-2002.
(BEIG-) EPIGENOMICS AG.
(ery Match 55.1%; Score 36.2; DB 6;
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Pred. No. 1.7e+02;
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Pred. No. 1.7e+02;
                                                                                                                                              Score 36.2; DB 6;
Pred. No. 1.7e+02;
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Pred. No. 1.7e+02;
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Pred. No. 1.7e+02;
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Pred. No. 1.7e+02;
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Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABLISZ693 standard; DNA; 6222 BP.
Human immune system associated gene SEQ ID NO:
WOZ002200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL32484 standard; DNA; 6244 BP.
Human immune system associated gene SEQ ID NO:
WO200200928-A2.
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(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.2;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%;
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10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%;
         5.1%;
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(EPIG-) EPIGENOMICS AG.
sry Match
5.1%;
st Local Similarity 44.9%;
                                                                                                             18-OCT-2001.

A (EPIG-) EPIGENOMICS AG.

Query Match

Best Local Similarity 54.0%;
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RESULT 1425
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       Query Match
Best Local Similarity
RESULT 1423
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ID ABL32693 standard;
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Best Local Similarity
RESULT 1431
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PD
PA
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DNA repair gene fragment complementary to#7
                                                                                       Drosophila melanogaster genomic polynucleotide SEQ ID NO 3265
WO200171042-A2,
27-SEP-2001.
                                                                                                                                                              Length 3296;
                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 5647;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 5647;
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                       Length 2396
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WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                        ABLJJS66 standard; DNA; 5647 BP.
Human immune system associated gene SEQ ID NO: 1539
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL70355 standard; DNA; 5647 BP.
Chemically treated cell signalling DNA sequence#123
WO200202807-A2.
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(POLY-) POLYMUN SCI INMUNOBIOLOGISCHE FORSCHUNG.

5.1%; Score 36.2; DB 6;

5t Local Similarity 51.6%; Pred. No. 1.4e+02;
                                                                                                                                                            5.1%; Score 36.2; DB 4; 45.5%; Pred. No. 1.5e+02;
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18-001.
(EPIG-2001.
(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.2; DB 6;
ery Match
50.3%; Pred. No. 1.7e+02;
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(UYMA) UNIV MACQUARIE.
1.17 Match
1.17 Match
1.17 Match 53.1%; Pred. No. 1.7e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                           5.1%; Score 36.2; DB 6; 50.3%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                     Score 36.2; DB 6;
Pred. No. 1.7e+02;
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Pred. No. 1.7e+02;
                                                                                                                                                                                                                              Human metastasis associated gene SEQ ID NO: 180 WO200177376-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAQ11710 standard; DNA; 5852 BP.
Dictyostellum plasmid Ddp2 containing Rep gene.
WO9106644-A.
                                                                       ABL17264 standard; DNA; 3296 BP.
                                                                                                                                                                                                              ABL34627 standard; DNA; 5539 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DNA; 5647 BP.
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03-JAN-2002
(EPIG-) EPIGENOMICS AG.
0TV MATCh | 5.1%;
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Chemically treated DNA r
WO200191622-A2.
01-NOV-2001.
(EPIG-) EPIGENOMICS AG.
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(BPIG-) BPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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                               Best Local Similarity RESULT 1414
                                                                                                                                                                           Best Local Similarity RESULT 1415
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RESULT 1416
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Best Local Similarity
RESULT 1420
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Best Local Similarity
RESULT 1419
                                                                                                                                            (PEKE ) PE CORP NY
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Query Match

Query Match

Query Match

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Best Local Similarity
RESULT 1450
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Tumour suppressor gene derived chemically modified sequence #231.
WO200168912-A2.
                                                                                                                                                                                              AAS46760 standard; DNA; 7900 BP.
Tumour suppressor gene derived chemically modified sequence #484.
WO200168912-A2.
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(EPIG-) EPIGENOMICS AG.
ery Match 5-1%; Score 36.2; DB 4; Length 12356;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.1%; Score 36.2; DB 6; Length 11944; 55.0%; Pred. No. 2e+02;
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                                                                                                                                                            Length 7589;
                                                                                                                                                                                                                                                                    Length 7900
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Human gene regulation-associated gene oligonucleotide #115.
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Signal transduction associated gene modified DNA #46.
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WO200200928-A2.
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Chemically treated cell signalling DNA sequence#48.
02020202807-A2.
10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                        ABL32880 standard; DNA; 10197 BP.
Human immune system associated gene SEQ ID NO: 853
W0200200928-A2.
(SPIG-) EPIGENOMICS AG.
                                                                                                    Human immune system associated gene SEQ ID NO: 921 W0200200928-A2. 03-JAN-2002. (BPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                       ABL32063 standard; DNA; 9905 BP.
Human immune system associated gene SEQ ID NO: 36
WC200200928-A2.
                                                                                                                                                             5.1%; Score 36.2; DB 6; 54.0%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 36.2; DB 6;
Pred. No. 1.9e+02;
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Pred. No. 1.9e+02;
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Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                      Score 36.2; DB 4;
Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 36.2; DB 6;
Pred. No. 1.9e+02;
                                                                54.0%; Pred. No. 1.8e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABK31248 standard; DNA; 10543 BP.
                                                                                          ABL32948 standard; DNA; 7589 BP
                                                                                                                                                                                                                                  WOZUGEZ-
20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
TTV MATCh '-11arity 46.9%;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%;
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity 5
RESULT 1432
ID ABL32948 stand-
DE Human immur
PN WO2020r
PD 03-7
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 1437
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                                                                                                                                                                        Best Local Similarity
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Best Local Similarity
RESULT 1434
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                                                                                                                                                                Query Match
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ABL70180 standard; DNA; 15923 BP.
Chemically treated cell signalling DNA sequence complementary to#35.
WO202202807-A2.
                                                                                                                                                                                                                                                                                                            ABK31221 standard; DNA; 15923 BP.
Signal transduction associated gene modified complementary DNA #32.
WO200200926-A2.
(BPIG-) EPIGENOMICS AG.
AAA46554 standard; DNA; 12986 BP.
Tumour suppressor gene derived chemically modified sequence #276.
Tumour Suppressor gene derived chemically modified sequence #276.
20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.2; DB 4; Length 12986; st Local Similarity 48.3%; Pred. No. 2e+02;
                                                                                                                                                                     Tumour suppressor gene derived chemically modified sequence #427. WO200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          NO 243
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 15923;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 17674;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human gene regulation-associated gene oligonucleotide #88
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human chemically modified disease associated gene SEQ ID WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABQ66981 standard; DNA; 23695 BP.

Human angiogenesis associated polynucleotide SEQ ID NO
WO200246454-A2.

13-UIN-2002.

(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human immune system associated gene SEQ ID NO: 1906
WO200200928-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL33344 standard; DNA; 17674 BP.
Human immune system associated gene SEQ ID NO: 1317
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 36.2; DB 6;
Pred. No. 2.3e+02;
                                                                                                                                                                                                         20-SEP-2001.
(BPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
(EPIG-) Score 36.2; DB 4;
(ery Match 50.3%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                           Score 36.2; DB 6;
Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 36.2; DB 6;
Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5.1%; Score 36.2; DB 6; 54.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 36.2; DB 6;
Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.1%; Score 36.2; DB 6; 49.2%; Pred. No: 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 36.2; DB 6;
Pred. No. 2.2e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABN80226 standard; DNA; 19236 BP.
                                                                                                                                                         AAS46704 standard; DNA; 14615 BP
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.1%;
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ry Match
t Local Similarity 54.0%;
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A (EPIG-) EPIGENOMICS AG.
Query Match 5
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity RESULT 1445
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RESULT 1448
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                                                                                                                 Best Local Similarity
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Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABK31511 standard; DNA; 47108 BP.
Signal transduction associated gene modified complementary DNA #177.
03-JAN-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 170245;
                                                                                                                                                                                                                                                                                             5.1%; Score 36.2; DB 12; Length 104245; 56.2%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 36.2; DB 12; Length 104245; Pred. No. 3.1e+02;
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                                                                                                                                                                                   Length 53552;
                                                                    Length 47108
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADP13586 standard; DNA; 170245 BP.
Renal cell carcinoma differentially expressed gene #322.
WO2004048933-A2.
                                                                                                                                                                                                                         ADG86768 standard; DNA; 104245 BP.
Human clone RP1-109F14 from chromosome 6p21.2-21.3.
US2003224514-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 36.2; DB 12;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%; Score 36.2; DB 12; 56.2%; Pred. No. 3.4e+02;
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Pred. No. 3.2e+02;
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Pred. No. 3.2e+02;
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Pred. No. 3.2e+02;
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Pred. No. 3.2e+02;
                                             03-JAN-ZUUG.
(EPIG-) EPIGENOMICS AG.
5.1%; Score 36.2; DB 6;
rry Match
rran cimilarity 48.3%; Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.1%; Score 36.2; DB 6; 50.3%; Pred. No. 3.2e+02;
                                                                                                                                                                 (WHED ) WHITEHEAD INST BIOMEDICAL RES.

ry Match
5.1%; Score 36.2; DB 5;
t Local Similarity 51.6%; Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 36.2; DB 2;
Pred. No. 3.2e+02;
                                                                                                 AMS13655 standard; DNA; 53552 BP.
Genomic DNA sequence encoding human hspG15.
WO200166752-A2.
                                                                                                                                                                                                                                                                                                                                      ADL34706 standard; DNA; 104245 BP.
Human PPAR-delta DNA fragment SEQ ID 4.
US2004063129-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA; 490885 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADK16049 standard; DNA; 49088
Naroarchaeum equitans genome.
WO200309344-A2.
13-NOV-2003.
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                                                                                                                                                                                                                                                                 04-DEC-2003.
(ISIS-) ISIS PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ᅜᇤ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (TWIN/) TWINE N C.
(BURC/) BURCZYNSKI M
(TREP/) TREPICCHIO W
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (DIVE-) DIVERSA CORP.
                                                                                                                                                                                                                                                                                                                                                                                01-APR-2004.
(GAAR/) GAARDE W.
(FREI/) FREIER S M.
(WATT/) WATT A T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (DORN/) DORNER A.
(STOV/) STOVER J.A.
(SLON/) SLONI D.K.
                                                                                                                                                                                           Best Local Similarity RESULT 1452
                                                                                              RESULT 1451
ID AAS13655 standard;
                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 1453
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 1461
                                                                     Query Match
Best Local Similarity
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Best Local Similarity
RESULT 1454
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Best Local Similarity
RESULT 1456
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Best Local Similarity
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Best Local Similarity
RESULT 1455
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Best Local Similarity
RESULT 1459
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Best Local Similarity
RESULT 1460
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RESULT 1457
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(AMHP ) WYETH.
                                                                                                                                                       13-SEP-2001
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ADAMBOUDS Standard; CDNA; 424 BP.
Bovine EST associated with lactation/muscle/fat deposition #11218.
US2002137139-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABX49849 standard; cDNA; 375 BP.
Bovine EST associated with lactation/muscle/fat deposition #15014.
US2002137139-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABX47746 standard; cDNA; 442 BP.
Wexine EST associated with lactation/muscle/fat deposition #12911.
US2002137139-A1.
26-SBP-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  LOLING BIOTECHNOLOGIE AG.

Best Local Similarity 56.2%; Pred. No. 3.5e+02;

RESULT 1466

ID ADB92119 standard; DNA; 189013 BP.

DE Human MRI related DNA sequence TP

PN W02003013535-A2.

PD 20-FEB-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 189013;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.1%; Score 36.2; DB 10; Length 189013; 56.2%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                   Length 189013;
                                                                                                                        Length 189013;
RESULT 1462

ID ACF62741 standard; DNA; 189013 BP.

ID Canner based on CYP3A5 related polynucleotide SEQ ID NO:669.

PN WO2003013534-A2.

PD 20-FEB-2003.

PD 20-FEB-2003 BIOTECHNOLOGIE AG.

PA (EPID-) BPIDAUROS BIOTECHNOLOGIE AG.

Query Match

Query Match

18; Score 36.2; DB 8; Length 1890
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 424;
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                                                                                                                                                                                                 MRP1 based cancer related nucleic acid SEQ ID NO:669.WO2003013533-A2.
                                                                                                                                                                                                                                                                                                                                                                                            GD-FEB-2003.

(EPID-) BPIDAUROS BIOTECHNOLOGIB AG.

ery Match
5.1%; Score 36.2; DB 10;

""""" Similarity 56.2%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                              WOZUUSALZONG.
20-FEB-2003.
(EPID) EPIDAUROS BIOTECHNOLOGIE AG.
(ERID) BATANATCh
5.1%; Score 36.2; DB 8;
--- cimilarity 56.2%; Pred. No. 3.5e+02;
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Pred. No. 1.1e+02;
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Pred. No. 1.1e+02;
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Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                       ADB87945 standard; DNA; 189013 BP.
Human UGT1A1 gene sequence SEQ ID NO:669
WO2003013536-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                20-FEB-2003.
(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                                 DNA; 189013 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.0%;
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(BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAOON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (BYAT/) BYATT J C. (MATH/) MATHIALAGAN N.
                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
RESULT 1465
                                                                                                                                                                                   ADB20856 standard;
                                                                                                                                                                                                                                                                                   Ouery Match
Best Local Similarity
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(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Local Similarity
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Length 2645;

Length 2729;

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ID AAH54890 standard; DNA; 3465 BP.

DB S. epidermidis genomic polynucleotide sequence SEQ ID NO:4254.

PN WO200134899-A2.

PD 17-MAY-2001.

PA (GLAX ) GLAXO GROUP LTD.

Query Match

S.0%; Score 36; DB 4; Length 3465;

Best Local Similarity 47.4%; Pred. No. 1.7e+02;

RESULT 1487

ID ACF62820 standard; DNA; 3664 BP.

DE Colon cancer analysis related genomic DNA SEQ ID NO:69.

PN WO2003014388-A2.

PD 20-FEB-2003.

PA (S.0%; Score 36; DB 8; Length 3664;

Query Match

S.0%; Score 36; DB 8; Length 3664;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DE AA4519 standard; DNA; 2865 BP.

DE Tumour suppressor gene derived chemically modified sequence #41.

DE Tumour suppressor gene derived chemically modified sequence #41.

DE 20-SEP-2001.

PA (EPIG-) EPIGENOMICS AG.

Query Match

Best Local Similarity 49.0%; Pred. No. 1.6e+02;

RESULT 1483

ID ABN800505 standard; DNA; 2865 BP.

DE Human chemically modified disease associated gene SEQ ID NO 67.

PN W02002002027-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     S. epidermidis genomic polynucleotide sequence SEQ ID NO:3503.
WO200134809-A2.
    5.0%; Score 36; DB 6; Length 2645; 52.7%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                 Length 2729;
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Best Local Similarity 49.0%; Score 36; DB 6; Len.

RESULT 1484

ID ARZ36285 standard; cDNA; 3196 BP.

BN WO200283876.A2.

PD 24-OCT-20^^
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            авизьс285 standard; cDNA; 3196 BP.
Human secretory polynucleotide SPTM SEQ ID NO 449.
W0200233876-A2.
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PA (INCY-) INCYTE GENOMICS INC.
Query Match
Best Local Similarity 49.5%; Pred. No. 1.7e+02;
RESULT 1485
                                                                                                                      USZUCZ-2003.
27-MAR-2003.
(HUMA-) HUMAN GENOME SCI INC.
1ery Match
52.7%; Score 36; DB 10;
52.7%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                    PD 07-FEB-2001.

PA (HELL-) HELIX RES INST.

Query Match 5.0%; Score 36; DB 4; 1
Best Local Similarity 49.5%; Pred. No. 1.6e+02;
RESULT 1481
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.0%; Score 36; DB 12; 49.5%; Pred. No. 1.6e+02;
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Pred. No. 1.7e+02;
                                                                                       Human cDNA from extracellular matrix gene 66.
US2003059875-A1.
                                                                                                                                                                                                                             AAH117823 standard; CDNA; 2729 BP.
Human CDNA sequence SEQ ID NO:17494.
EP1074617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                         ADO20361 standard; cDNA; 2729 BP. Human PRO polymucleotide #628. BP. 27-MAY-2004. GENENTECH INC.
                                                              ADC10608 standard; cDNA; 2645 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAH54139 standard; DNA; 3335 BP.
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(GLAX ) GLAXO GROUP LTD.
(GLAX ) GLAXO 47.0%;
Query Match
Best Local Similarity
RESULT 1479
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Best Local Similarity
RESULT 1480
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RESULT 1482
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Best Local Similarity
RESULT 1486
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AA255560 standard; cDNA; 2262 BP.
Human secreted protein clone ye7_1 nucleotide sequence SEQ ID NO:171.
WO9958642-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADE37766 standard; DNA; 1984 BP.
Human chemically treated calcitonin nucleotide sequence SEQ ID NO:12.
W02003072820-A2.
04-SEP-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 1984;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Chemically treated cell signalling DNA sequence#252.
Chemically treated cell signalling DNA sequence#252.
WOZD0202807-A2.
10-JAN-2002.
10-JAN-2002.
EPIGENOMICS AG.
5.0%; Score 36; DB 6; Length 1984; ery Match
St. Local Similarity 50.6%; Pred. No. 1.5e+02;
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(HUMA-) HUMAN GENOME SCI INC.
6ry Match
5.0%; Score 36; DB 4; Length 2645;
7.7.1 Similarity 52.7%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5.0%; Score 36; DB 6; Length 1984; 50.6%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 2262;
                      ADL44075 standard; DNA; 469 BP.

Human ovarian cancer DNA marker #17965.

WAZ00170979-A2.

27-SEP-2001.

MILLENNIUM PREDICTIVE MEDICINE INC.

Guery Match

5.0%; Score 36; DB 5; Length 469;
Best Local Similarity 55.2%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                               Length 480;
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LUNA; 2262 BP.

FU 18-NOV-1999.

PA (GEMY) GENETICS INST INC.

QUERY MATCH

RESULT 1477

ID AAS31262 standard; CDNA; 2645 BP.

DE Human CDNA encoding a novel ...

PD 02-AUG-2001

PD 02-AUG-2001
                                                                                                                                                                                                                    Human cancer related polynucleotide SEQ ID NO 3977
WO200214500-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Molocology, 04-SEP-2003, (EPIG-) EPIGENOMICS AG, 5.0%; Score 36; DB 10; ery Match 7.1.5e+02; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                   ABL34634 standard; DNA; 1984 BP.
Human metastasis associated gene SEQ ID NO: 187.
NO20017376-A2.
18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                             Score 36; DB 6;
Pred. No. 1.1e+02;
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Local Similarity 50.6%; Pred. No. 1.5e+02;
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Human polynuclectide SEQ ID NO 76.
US2002042386-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADB54175 standard; DNA; 1984 BP.
Pretreated genomic DNA region 99.
WO2003072821-A2.
                                                                                                                                                                                                ABN64010 standard; cDNA; 480 BP.
                                                                                                                                                                                                                                                                                                                             5.0%;
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(CHIR ) CHIRON CORP.
(HYSE-) HYSEQ INC.
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Best Local Similarity
RESULT 1473
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(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
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RESULT 1478
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RESULT 1472
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      RESULT 1470
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43.1%; Pred. No. 1.7e+02;

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Haematopoietic cell proliferation disorder related DNA sequence #268.
WO200277272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABQ75107 standard; cDNA; 4985 BP.
Anopheles gambiae odourant receptor 2 genomic DNA SEQ ID NO:10.
WO200259274-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 4985;
                                                                                              03-0CT-2002.
(BPIG-) EPIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
5.0%; Score 36; DB 8; Length 3664;
lery Match
5.0%; Pred. No. 1.7e+02;
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WO200200928-A2.
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(EPG-0. EPIGENOMICS AG.

(EPT Match 5.0%; Score 36; DB 6; I

5.0%; Score 36; DB 6; I

5.0%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                            23-AUG-2001.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

ery Match

5.0%; Score 36; DB 5; I

ref 1003 Similarity 49.5%; Pred. No. 1.8e+02;
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sry Match 5.0%; Score 36; DB 5; I

t Local Similarity 49.5%; Pred. No. 1.8e+02;
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60.0%; Pred. No. 1.8e+02;
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Human immune system associated gene SEQ ID NO:
WO200200928-A2.
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Pred. No. 1.9e+02;
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Best Local Similarity 49.5%; Pred. No. 1.8e+
RESULT 1490
DD ABV21NO standard; CDNA; 4085 BP.
DB Human prostate expression marker cDNA 21771.
PN WO200160860-A2.
PD MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                       Human prostate expression marker cDNA 27594.
WC200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ACF79720 standard; DNA; 4985 BP.
Mosquito odorant receptor 2 genomic DNA.
WO2003076590-A2.
                                                                                                                                                                                                       CDNA; 4085 BP
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                                               ABZ10128 standard; DNA; 3664 BP
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.0%;
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(UYVA-) UNIV VANDERBILT.
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(EPIG-) EPIGENOMICS AG.
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RESULT 1491
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Best Local Similarity
RESULT 1495
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Best Local Similarity
RESULT 1496
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Best Local Similarity
RESULT 1488
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RESULT 1489
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RESULT 1492
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RESULT 1497

ID ABL70594 standard; DNA; 6061 BP.

ID Chemically treated cell signalling DNA sequence complementary to#242.

PD Chemically treated cell signalling DNA sequence complementary to#242.

PD 10-JAN-2002.

PA (EPIG-) EPIGENOMICS AG.

Query Match

5.0%; Score 36; DB 6; Length 6061;

Best Local Similarity 50.6%; Pred. No. 1.9e+02;

RESULT 1498
                                                                                                                                                                                                                                                                                                                                                                                                                                              ID NO 106.
                                                                                                                                                                                                                          DE Human gene regulation-associated gene oligonucleotide #266.
PN WO200177375-A2.
PD 18-OCT-2001.
PA (EPIG-) EPIGENOMICS AG.
Query Match
5.0%; Score 36; DB 6; Length 6061;
Best Local Similarity 50.6%; Pred. No. 1.9e+02;
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Human immune system associated gene SEQ ID NO: 793.
WO200200928-A2.
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Pred. No. 1.9e+02;
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Pred. No. 1.9e+02;
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PA (EPIG-) EPIGENOMICS AG.

Query Match 5.0%;

Best Local Similarity 51.2%;

RESULT 1500

ID ABL32820 standard; DNA; 6123
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(EPIG-) EPIGENOMICS AG.
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Sequence 11, Appl
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AG311121 AV326659 2 BE086569 2 BE086569 2 BE724994 9 CE646245 6 CD68251 4 BM522061 6 CD68251 6 CD85593 6 CD85594 6 CD85594 6 CD85594 6 CD85594	ALIGNMENTS 663 bp mRNA ens Mast cell library 519, mRNA sequence. 23690	mates; imates; id Oya,Y id Oya,Y ib, URL: lfiers in sapie	imilarity 95.3%; Score 527.8; DB 1; Conservative 0; Mismatches 23; ATATATCATTATCATTAATCATAATGHEN TCTTTTTTTGGATTTTTTTTTTTTTTTTTTTTTTTTTTT
2020 2020 2020 2020 2020 2020 2020 202	no sapiens M 302302-019, GI:14623690 s (human)	**Metazoa; Chorresta; Prime Eutheria; Prime 1 to 663) **Oshida,T. and A sequencing A sequencing A sequencing A sequencing A sequencing and 1201) **Yuji Sugita and 1797-2281 **A-797-2281 **A-797-2282 **Uji@genox.co.jp **Location/Qualif.Cocation/Qualif.Type="MRNA," **Morl Erype="MRNA," **Morl Erype="Morl Erype="Mass." **Morl Erype="Mass." **M	74.0%; 95.3%; 95.3%; ative TAITTAIC TAITINAIC TAITINAIC NAAITTIC AAGCTAIT AAGAAAIAA
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	CA970412 CCLL03a08 BH331915 CH230-46N CC483944 CH240-312 AJ599212 Arabidops BM395523 50072-2-9 AL25470 Tetraodon CL835279 OR CBa006 AZ684844 ENTIU33TF CG030862 PUTFP83TB AG494504 Mus muscu AL211196 Tetraodon AL070621 Drosophil B10881 F2446-5p6.1 CL475079 SAIL 231	AG388848 Mus muscu AL051882 Drosophil AL104701 Drosophil AG38792 AGENCOURT AG38792 Mus muscu CC239214 CH261-127 BI490178 603031949 CL077343 CH216-150 CL077343 CH216-144 CL461494 SALL 1148 AL104520 Drosophil AL07523 Drosophil AL06896 Drosophil AL106896 Drosophil AL106898 Drosophil CR591130 full-leng BM917623 AGENCOURT AG310974 Mus muscu AG719752 HS S572 B RX40765 BX40066	BG519032 602577955 CL491661 SAIL 559 BU165262 AGENCOURT AL067050 Drosophil AL106262 Drosophil AL106455 Mus muscu AL106476 Drosophil BM547485 AGENCOURT CL019707 CH216-5M2 AG280144 Mus muscu CF22265 AGENCOURT CL019709 CH216-5M2 AG371391 Mus muscu CF22265 AGENCOURT AG347035 Mus muscu CF22265 AGENCOURT AG34035 Mus muscu CG56595 AGENCOURT AG349990 Mus muscu CG754506 P049-4-H0 AG280271 Mus muscu CG754506 P049-4-H0 AG280125 Mus muscu AG280271 Mus muscu AG280271 Mus muscu AG280271 Mus muscu AG280275 Mus muscu
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Akinret.B., Nierman, W., Feldblyum, T., Malek, J., Shatsman, S., Akinret.B., Levins, M., Megann, S., Tsegaye, G., Geer, K., Krol, M., de Jong, P. and Fraser.C.M.

Nouse BAC End Sequences from Library RPCI-23

AL Unpublished (1999)

Other GSS: RPCI-23-30A15.TJ

Contact: Shaying Zhao

Opatrment of Eukaryotic Genomics

The Institute for Genomic Research

9712 Medical Center Dr., Rockville, MD 20850, USA

Tel: 301 838 0200

Fax: 301 838 0200

Sequences (http://www.igr.org/tdb/bac_ends/mouse/bac_end_intro.html

Plate: 30 row: A column: 15

Seq primer: T7

Class: BAC ends
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 RPCI-23-30A15.TV RPCI-23 Mus musculus genomic clone RPCI-23-30A15, genomic survey sequence.
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Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 663)
kb (range 0.60-3.5 kb). 15/15 colonies contained inserts by PCR. This library was enriched for full-length clones and was constructed by Clontech Laboratories (Palo Alto, CA). Note: this is a NIH_MGC Library."
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                                                                                                                                               Score 268.2; DB 6,
Pred. No. 2.5e-50;
0; Mismatches 3;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               TGTGACTCCTACCAAAGCTGTCAAAAACAACGG 523
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                                                                                                                                                  37.6%;
98.9%;
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Best Local Similarity 98.9
Matches 270; Conservative
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/clone="Infage:30352338"
/lab host="DH108 (T1 phage-resistant)"
/clone lib="NIH MGC 184
/clone lib="NIH MGC 184
/note="Organ: Pooled-Glandular; Vector: pDNR-LIB; Site_1:
Sfil (ggccattatggcc); Site_2: Sfil (ggccgcctcggcc);
Library is oligo-dT primed and directionally cloned. cDNR
was prepared from a glandular pool of tissues from thyoid,
parathyroid, adrenal, cortex and pineal gland. 5' and 3'
adaptors were used in cloning as follows: 5' adaptor
sequence: 5'-CACGGCCARTATGGCC-3' and 3' adaptor sequence:
5'-ATTCTAGAGGCCGAGGCGACATG-dT(30)BN-3' (where B = A,
C, or G and N = A, C, G, or T). Average insert size 1.38
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Email: cgapbs-remail.nih.gov
Tissue Procurement: Dr. Michael Brownstein and Dr. Miklos Palkovits
cDNA Library Preparation: CLOYTECH Laboratories, Inc.
cDNA Library Arrayed by: The I.M.A.G.E. Consortium (ILNL)
DNA Sequencing by: Agencourt Bioscience Corporation
Clone distribution: MGC clone distribution information can be
found through the I.M.A.G.E. Consortium/LLNL at:
Plate: NDCM151 row: e column: 19
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Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 800)
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                                        TGACTCAAGAGGGTTAATTCTTGGTGCTGAAGCCTGGGGGCAGGGGTGTAAAGAAAAACAC
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              NIH-MGC http://mgc.nci.nih.gov/.
National Institutes of Health, Mammalian Gene Collection (MGC)
Unpublished (1999)
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AGENCOURT 13785021 NIH MGC 184 Homo sapiens cDNA clone
IMAGE:30352338 5', mRNA sequence.
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/db_xref="taxon:9606"
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Homo sapiens

Bukaryota, Metazoa, Chordata, Craniata, Vertebrata; Buteleostomi,

Bukaryota, Metazoa, Chordata, Craniata, Vertebrata; Buteleostomi,

(bases 1 to 673)

Gu,Y., Peng,Y., Song,H., Huang,Q., Yang,Y., Gao,G., Xiao,H., Xu,X.,

Gu,Y., Peng,Y., Song,H., Huang,Q., Yang,Y., Gao,G., Xiao,H., Xu,X.,

Li,N., Oian,B., Liu,F., Qu,J., Gao,X., Cheng,Z., Xu,Z., Zeng,L.,

Xu,S., Gu,W., Tu,Y., Jia,J., Fu,G., Ren,S., Zhong,M., Lu,G., Hu,R.,

Chen,Z., and Han,Z.

Unpublished (2000)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AV721179 HTB Homo sapiens cDNA clone HTBAKF11 5', mRNA sequence. AV721179
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Chinese National Human Genome Center at Shanghai
351 Guo Shoujing Road, Zhangjiang Hi-Tech Park, Pudong, Shanghai
201203, P. R. China
Tel: 86-21-50801919 (ex.45)
Fax: 86-21-50801920 sh.cn
Email: hanzg@chgc.sh.cn
This clone is available at CHGC in Shanghai.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    /tissue_type="Hypothalamus"
/dev_stage="Adult"
/lab_host="SOLR"
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1. 673
1. 673
1. 673
1. mol = "Homo sapiens"
/mol = "Type="mRRA"
/db_xref="taxon:9606"
                                                                                                                                                                                                                                                                                                                                                                                                                             stringency conditions."
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       452 AAAGCTGTCAAAACCACAGG 433
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Best Local Similarity 99.3%;
Matches 139; Conservative
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JOURNAL
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AUTHORS
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/lab host="DHIDB"
/clone lib="RPCI-23"
/clone lib="RPCI-23"
/note="Uorgan: Kidney/Brain; Vector: pBACe3.6; Site_1:
ECGRI; Site 2: ECGRI; Female G57BL/6J mouse kidney and/or
brain genomic DNA was isolated and partially digested
with a combination of EcoRI and EcoRI Methylase. Size
selected DNA was cloned into the pBACe3.6 vector at the
ECGRI sites. The ligation products were transformed into
DHIOB electrocompetent cells (BRL Life Technologies). "
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Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
1 (bases I to 659)
Dias Neto, E., Garcia Correa, R., Verjovski-Almeida, S., Briones, M.R.,
Nagai, M.A., da Silva, W. Jr., Zago, M.A., Bordin, S., Costa, F.F.,
Goldman, G.H., Carvalho, A.F., Matsukuma, A., Baia, G.S., Simpson, D.H.,
Brunstein, A., deoliveira, P.S., Bucher, P., Jongeneel, C.V.,
O'Hare, M.J., Soares, F., Brentani, R.R., Reis, L.F., de Souza, S.J. and
                                                                                                                                                                                                                                                                                                                                  4
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IL3-MT0267-050101-437-A02 MT0267 Homo sapiens cDNA, mRNA sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    231 GTTCGTGACGTATGAAGCATTTTCGTGACAATTCAGGGAGAAATCCAGAGGAAAAA 172
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                CTACTITCIATCAAGAAAIAAAGAGAACCACAGTCAAACCACACACAATCATCTITAGAAGA 237
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This sequence was derived from the FAPESF/LICR Human Cancer Genome
                                                                                                                                                                                                                                                                                                                                                                                                                          351 AACATACAGTTTATTTATCAATAACCATAGGCATCCCCTATATATGTCCCATAAATATGA 292
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                60 GITTTGGGATTTTAATTTTCAAACAGCAGAATGACAI--TTTTTCTGTCACTATTATT 117
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 232
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              171 CAACTTCCTATCAAGAAATAAAGAGAACCACA---AGCCCACAGAATCATCTTAGATGA 115
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               238 CAGIGIGACICCIACCAAAGCIGICAAAACCACAGGGCAAGGGCAIAGITAAAGGA---- 292
                                                                                                                                                                                                                                                                                                                                                                              1 AATATATCATCTATTTATCATTAATCAATAATGTATTCT-TTTATTCCAATAACATTTGG 59
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Laboratory of Cancer Genetics
Ludwig Institute for Cancer Research
Rua Prof. Antonio Prudente 109, 4 andar, 01509-010, Sao Paulo-SP,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Shotgun sequencing of the human transcriptome with ORF expressed
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11; Gaps
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20202663
                                                                                                                                                                                                                                                                               Length 663;
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                                                                                                                                                                                                                                                                       Score 147.6; DB 8;
Pred. No. 5.6e-23;
0; Mismatches 89;
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Best Local Similarity 71.6%;
Matches 252; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Fax: +55-11-2707001
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Best Local Similarity
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Project. This entry can be seen in the following URL (http://www.ludwig.org.br/scripts/gethtml2.pl?tl=IL3&t2=IL3-MT0267-0501014-37-A02&t8=2501-01-05&t4=1)
Seq primer: puc 18 forward Seq primer: puc 18 forward Seq primer: puc 18 loward Location/Quality sequence stop:
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Pred. No. 6.8e-21;
0; Mismatches 1; Indels 0
                                                                                                                                                  organism="Homo sapiens"
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Mus musculus
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KEYWORDS
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                                                                              RESULT 7
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Smith, T.P.L., Roberts, A.J., Echternkamp, S.E., Chitko-McKown, C.G.,
Wray, J.E. and Keele, J.W.
A second set of bovine ESTs from pooled-tissue normalized libraries
Unpublished (2003)
Contact: Smith TPL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Email: smith@email.marc.usda.gov
Single pass sequencing. Bases called with phred v0.020425.c and
trimmed with the aid of the trim_alt option. Vector identified with
cross_match v0.990329.
Plate: FQY8014 row: B column: 7
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                                                                                                                                                                                                                                                                                                                                                                                          EST 25-MAR-2003
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                                                                                                                                                                                                                                                        214 ACCCACACAATCATCTTTAGAAGACAGTGTGACCTCCTACCAAAGCTGTCAAAACCACAGG 273
                                                                                                                                                                                                                                                                                 61 ACCCACACATCATCTTTAGAAGACAGTGTGACTCCTACCAAGCTGTCAAAAACCACAGG 120
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                                                                                                                                                                                                                    9
/clone_lib="HTB"
/note="Vector: pBluescript sk(-); Site_1: EcoRI; Site_2:
XhoI"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Cetartiodactyla, Ruminantia, Pecora, Bovidae,
Bovinae, Bos.
                                                                                                                                                                                                  1 AGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAGTCA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     /note="Vector: pcDNA3.1; Site 1: EcoRI; Site 2: NotI; Library made with RNA pooled from multiple tissues including liver, lung, hypothalamus, pituitary, and placenta/endometrium."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   134 GCTATTTGGAGATCCAATTCAGGAAGCAACACTTGGAGAATGGCTACTTTCTATCAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        244 GGTATTTGGAGATCCAGTTCAGGGAACAACTGTTGAAGAGTGACAGCTTTCCATCAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  194 AATAAAGAGAACCACAGTCAACCCACACAATCATCTTTAGAAGACAGTGTGACTCTACC
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                                                                                               Length 673;
                                                                                                                                                                                                                                                                                                                                                                                          CB420818 582 bp mRNA linear
593806 MARC 6BOV Bos taurus cDNA 5', mRNA sequence.
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                                                                                                                                     0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   USDA, ARS, US Meat Animal Research Center
PO Box 166, Clay Center, NE 68933-0166, USA
TEL: 402 762 4360
Fax: 402 762 4390
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      14.7%; Score 104.8; DB 6;
84.3%; Pred. No. 2.8e-13;
live 0; Mismatches 22;
                                                                                               DB 1;
1e-16;
                                                                                               16.8%; Score 120; DB
100.0%; Pred. No. 1e-
ive 0; Mismatches
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/db_xref="taxon:9913"
/tissue_type="pooled"
/lab_host="DH10B"
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Best Local Similarity 84.3%;
Matches 118; Conservative
                                                                                               Query Match
Best Local Similarity 100.
Matches 120; Conservative
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ORGANISM
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JOURNAL
COMMENT
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CB420818
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364 AGGCTCTACGACCACGG 383

S64 AGGCTCTACGACCACACGG 383

TITIO 0117544 RIKEN [tull=langth enriched be derived coll in each sequence.

S10 BY347544 RIKEN [tull=langth enriched] Be derived coll in each sequence.

BY35 BY37544 IG 126681923

BY375441 G126681923

BY375441 G126681923

BY375441 G126681923

BY375441 G12668192

BY375441 G12668192

BY3754741 G12668192

BY375471 G12668192

BY375471 G12668192

BY375471 G12668192

BY37547 G126681 G1268192

BY37547 G12681 G1268192

BY37547 G12681 G1268192

BY37547 G12681 G1
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Best Local Similarity 69.8
Matches 132; Conservative
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Nikaido, I., Osato, N., Kasukawa, T., Adachi, J., Bono, H., Kondo, S., Nikaido, I., Osato, N., Saito, R., Suzuki, H., Yamanaka, I., Kiyosawa, H., Yagi, K., Tomaru, Y., Hasegawa, Y., Nogami, A., Schonbach, C., Gojobori, T., Baldarelli, R., Hill, D.P., Bult, C., Hume, D.A., Quackenbush, J., Schriml, L.M., Kanapin, B., Marsuda, H., Batalov, S., Beisel, K.W., Blake, J.A., Bradt, D., Brusic, V., Chotani, L.E., Cousine, S., Dalla, E., Dragani, T.A., Fletcher, C.F., Forrest, A., Frazer, K.S., Gaasterland, T., Gariboldi, M., Gissi, C., Godzik, A., Gough, J., Grimmond, S., Gustincich, S., Hirokawa, N., Jackson, I.J., Jarvis, B.D., Kanai, A., Kawasawa, Y., Kedzierski, R.M., King, B.L., Konagaya, A., Kurochkin, I.V., Lee, Y., Lenhard, B., Lyons, P.A., Maglott, D.R., Maltais, L., Marchionni, L., McKenzie, L., Miki, H., Nagashima, T., Numata, K., Okido, T., Pavan, W.J., Pertea, G., Pesole, G.,
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nonredundant cDNA library. Genome Res. 11 (2), 281-289 (2001) CDNA library was prepared and sequenced in Mouse Genome Experienced for Research Group in Riken Genomic Sciences Center and Genome Science Laboratory in RIKEN. Division of Experimental Animal Research in Riken contributed to
                                                                                                                                                                         Tissues were provided by Dr. John Todd (Dept. of Medical Genetics Wellcome Trust Centre for Molecular Mechanisms in Disease Wellcome Trust/MRC building Addenbrookes Hospital Cambridge) whose
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               295 GTTCGTGACGTATGAAGCATTTTGGCGACACAATTCAGGGAGAAATCCAGGGAGAAAGA 354
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  60 GTTTTGGGATTTTAATTTTCAAACACAGCAGAATGACAT--TTTTTCTGTCACTATTATT 117
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                118 ATTGTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGG 177
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Mammalia, Butheria, Rodentia, Sciurognathi, Muridae, Murinae, Mus.
1 (bases 1 to 419)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    /cell_type="B6-derived CD11 +ve dendritic cells"
/clone_lib="RIKEN full-length enriched, B6-derived CD11
+ve dendritic cells"
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                                                                                                                                                                                                                                                                  assistance we gratefully acknowledge.
Please visit our web site (http://genome.gsc.riken.go.jp) for further details.
Location/Qualifiers
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           3;
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Pred. No. 6.8e-07;
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                                                                                                                                                                                                                                                                                                                                                                                                           /organism="Mus musculus"
/mol_type="mRNA"
/strain="C57BL/6J"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /db_xref="taxon:10090"
/clone="F730119008"
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69.8%;
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Best Local Similarity
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BY536666
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Petrovsky, N., Pillai, R., Pontius, J. U., Qi, D., Ramachandran, S., Ravaai, T., Reed, J. C., Reed, D. J., Reid, J. Ring, B. Z., Ringwald, M., Sandellin, A., Schneider, C., Semple, C.A., Setou, M., Shimada, K., Sultana, R., Takenaka, Y., Taylor, M., Stimada, K., Verardo, R., Wagner, L., Wahlestedt, C., Wang, Y., Watanabe, Y., Welse, C., Wilming, L.G., Wynshaw-Boris, A., Yanagisawa, M., Yang, I., Yang, L., Yan, Z., Zavolan, M., Zhu, Y., Zimmer, A., Carninci, P., Hayatsune, N., Birozane-Kishikawa, T., Konno, H., Nakamura, M., Sakaume, N., Saro, K., Shiraki, T., Waki, K., Kawai, J., Aizawa, K., Arakawa, T., Fukuda, S., Hara, A., Hashizume, W., Inotani, R., Ishii, Y., Itoh, M., Xagawa, I., Miyazaki, A., Saski, W., Sasaki, D., Shibata, K., Shinagawa, A., Yasunishi, A., Yoshino, M., Waterston, R., Lander, E.S., Analysis of the mouse transcriptome based on functional annotation of 60,770 full-length cDNs.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Email: genome-resignsc.riken.jp, URL:http://genome.gsc.riken.jp/
Aizawa,K., Akimura,T., Arakawa,T., Carninci,P., Fukuda,S.,
Hirozane,T., Inotani,K., Ishii,Y., Itoh,M., Xawai,U., Konno,H.,
Miyazaki,A., Murata,M., Nakamura,M., Nomura,K., Numazaki,R.,
Ohno,M., Sakai,K., Sakazume,N., Sasaki,D., Sato,K., Shibata,K.,
Shiraki,T., Tagami,M., Waki,K., Watahiki,A., Muramatsu,M. and
Hayashizaki,Y. Direct Submission
Computational Analysis of Full-Length Mouse cDNAs Compared with
Human Genome Sequences Mamm. Genome. 12, 673-677 (2001)
Normalization and subtraction of cap-trapper-selected cDNAs to
prepare full-length cDNA libraries for rapid discovery of new
genes. Genome Res. 10 (10), 1617-1650 (2000)
RIKEN integrated sequence analysis (RISA) system-184-format
sequencing pipeline with 384 multicapillary sequencer. Genome Res.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Laboratory for Genome Exploration Research Group, RIKEN Genomic Sciences Center(GSC), Yokohama Institute
The Institute of Physical and Chemical Research (RIKEN)
1-7-22 Suehiro-cho, Tsurumi-ku, Yokohama, Kanagawa 230-0045, Japan Tel: 81-45-503-9226
Fax: 81-45-503-9216
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Tissues were provided by Dr. John Todd (Dept. of Medical Genetics
Wellcome Trust Centre for Molecular Mechanisms in Disease Wellcome
Trust/MRC building Addenbrookes Hospital Cambridge) whose
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/clone_lib="RIXEN full-length enriched, NOD-derived CD11c
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Please visit our web site (http://genome.gsc.riken.go.jp) for further details.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Contact: Yoshihide Hayashizaki
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/strain="NOD"
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Direct Submission

Submitted (102-JUN-1999) Genoscope - Centre National de Sequencage :
Bp 191 91006 EVRY cedex - FRANCE (E-mail : seqref@genoscope.cns.fr

- Web : www.genoscope.cns.fr)

Determination of this BAC-end sequence was carried out as part of collaboration with the Berkeley Drosophila Genome Project (BDGP).

The BDGP is constructing a physical map of the Drosophila melanogaster genome using these BACs. For further information please see http://www.fruitfly.org The BDGP Drosophila melanogaster BAC library was prepared by Kazutoyo Osoegawa and Aaron Mammoser in Pieter de Jong's laboratory in the Department of Cancer Genetics at the Roswell Park Cancer Institute in Buffalo, NY. The library is named RPCI-9 and was constructed by partial EcoxI digestion of Drosophila DNA provided by the BDGP from the isogenic strain v2; cn bw sp, the same strain used for the BDGP's pl and BST libraries. A more detailed description of the library and how to order individual BAC clones, the entire library, or filters for hybridization from the BACPAC Resource Center can be found at http://www.fruitflalo.edu/drosophila_bac.htm.
                                                                                                                     CNS0039G 1101 bp DNA linear GSS 03-JUN-1999 Drosophila melanogaster genome survey sequence TET3 end of BAC #BACR08K10 of RPCI-98 library from Drosophila melanogaster (fruit
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     753
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   313
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                194 AATAAAGAGAACCACACTCAACCCACAATCATCTTTAGAAGACAGTGTGACTCCTACC 253
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        989 DAATDADGAGRRDGGRRRKDRKDRKDGDDDKKGGKKKKAAKKAAKWATKWMDDWDKDWK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        74 ATTTTCAAACACAGCAGAATGACATTTTTTTCTGTCACTATTATTATTGTTGGTATGTGAA
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                                                                                                                                                                                                                                                                                                                                                     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
1 (bases 1 to 110p)
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22; Conservative 251; Mismatches 277; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                /organism="Drosophila melanogaster"
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18.7%; Pred. No. 0.0089;
                                                                                                                                                                                                                                                                                                        Drosophila melanogaster (fruit fly)
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/db_xref="taxon:7227"
/clone="BACR08K10"
                                                                                                                                                                                             fly), genomic survey sequence.
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427 CAAGGCATCCCAAACTACAGG 447
                                                                                                                                                                                                                                                                                                                                  Drosophila melanogaster
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Best Local Simil
Matches 122; C
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602891529F1 NCI_CGAP_Lu29 Mus musculus cDNA clone IMAGE:5036647 5',
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                                                                                                                                                                        351 GTTCGTGACGTATGAAGCATTTTGGCGACACAATTCAGGGAGAAATCCAGAGGAGAAAGA 410
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Butheria, Rodentia, Sciurognathi, Muridae, Murinae, Mus.
1 (bases 1 to 855)
NIH-MGC http://mgc.nci.nih.gov/.
National Institutes of Health, Mammalian Gene Collection (MGC)
Unpublished (1999)
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/note="bggan: lung; Vector: pCMV-SPORT6; Site_1: Sall;
Site_2: Not1; Cloned unidirectionally. Primer: Oligo dT.
Library constructed by Life Technologies. Investigator
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/clone="IMAGE:5036647"
/tissue type="spontaneous tumor, metastatic to mammary.
Stem coll origin."
/lab_host="DH108"
                                                        118 ATTGTTGGTATGTGAGCTATTTGGAGATCCAATTCAGGAAGCAACACACATTGGAGAATGG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             pe.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Email: cgapbs-remail.nih.gov
Tissue Procurement: Gilbert Smith, Ph.D.
cDNA Library Preparation: Life Technologies, Inc.
cDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MGC clone distribution information can found through the I.M.A.G.E. Consortium/LLNL at:
http://mage.llh.gov
Plate: LLAM11101 row: j column: 08
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           providing samples: Gilbert Smith, NIH
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Contact: Robert Strausberg, Ph.D.
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High quality sequence stop: 825.
Location/Qualifiers
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/strain="CZECH II"
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CNSOOFUH
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           UNSUDUKY
T7 end of clone AWOAA009H09 of library AWOAA from strain CLIB 89 of ALA11257
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Submitted (07-SEP-2000) Genoscope - Centre National de Sequencage, 2 rue Gaston Cremieux, CP 5706, 91057 EVRY cedex, FRANCE. (E-mail: seqref@genoscope.ons.fr - Web: www.genoscope.ons.fr)
This GSS is part of a random genomic sequencing program of thirteen yeast species: Saccharomyces bayanus var. uvarum, Saccharomyces
                                                                                                                                                         493
                                                                                                                                                                                                     573
                                                                                                                                                                                                                                               553
                                                                                                                                                                                                                                                                                           513
                                                                                                                                                                                                                                                                                                                                     554 GGAGATCATATAATTGATACAAATAAAAGAAAAGTGTTCTCCCCTTACAGAATTGAC 613
                                                                                                                                                                                                                                                                                                                                                              Souciet, J.L., Aigle, M., Artiguenave, F., Blandin, G., Bolotin-Fukuhara, M., Bon, E., Brottier, P., Casaregola, S., Bolotin-Fukuhara, M., Bon, E., Brottier, P., Casaregola, S., Bolotin-Pukup, Duljon, B., Durenes, P., Lepingle, A., Llorete, B., Malpertuy, A., Neuveglise, C., Ozier-Kalogoropoulos, O., Potier, S. Saurin, W., Tekaia, F., Toffano-Nioche, C., Wesolowski-Louvel, M., Wincker, P. and Weissenbach, J. Genomic exploration of the hemiascomycetous yeasts: 1. A set of yeast species for molecular evolution studies
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Casaregola, S., Neuveglise, C., Lepingle, A., Bon, E., Feynerol, C., Artiguenave, F., Wincker, P. and Gaillardin, C. Genomic exploration of the hemiascomycetous yeasts: 17. Yarrowia
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Yarrowia lipolytica
Yarrowia lipolytica
Eukaryota; Fungi, Ascomycota; Saccharomycotina; Saccharomycetes;
Saccharomycetales; Dipodascaceae; Yarrowia.
1 (bases 1 to 759)
| .... | .... | .... | .... | .... | TWDRDDDDARDDBRRRRGDDGARARKRGDDGARARKRGDGARAGK TWKRRRRRRDTRWDDADADDARDDRRRRGDDGADAGK
                                                                   374 TTGTAAATTTAAGGCAAATACACATATTAGTATTACCTTAGTGTAATGTATCCCTGTCAT
                                                                                                                                                                                                   632 WDWWTWKAWDWAKWDWKTRADRWDRWAADTWTDARKADRDWAKARAWRARRDRARAD
                                                                                                                                                                                                                                                                                           434 ATATACAATAAGGTGAAATTATAAGTACCCTATGCAGTTGGCTGGACAGTTCTAAATTGG
                                                                                                                                                                                                                                                 494 ACTITATTAATTTTTAAAATCAGTAACTGATTTATCACTGGCTATGTGCTTAGATCTACA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       614 ATTITAAATGCGATACAGTTAGAATAGGAAATATGACATTAGAAAGGAAGAAT
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Direct Submission

Submitted (02-UTM-1999) Genoscope - Centre National de Sequencage:
BP 191 91006 EVRY cedex - FRANCE (E-mail: seqref@genoscope.cns.fr
- Web: www.genoscope.cns.fr)

Determination of this BAC-end sequence was carried out as part of a collaboration with the Berkeley Drosophila Genome Project (BDGP).

The BDGP is constructing a physical map of the Drosophila melanogaster genome using these BACs. For further information please see http://www.fruitfly.org The BDGP Drosophila melanogaster BAC library was prepared by Kazutoyo Geogawa and Aaron Mammoser in Pieter de Jong's laboratory in the Department of Cancer Genetics at the Roswell Park Cancer Institute in Buffalo, NY. The library is named RPCI-98 and was constructed by partial ECORI digestion of Drosophila DNA provided by the BDGP from the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  linear GSS 03-JUN-1999
nce TET3 end of BAC:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       647 TGACATTAGAAAGGAAGAATGACAGGAGAAAGGAAAGAAGGAAAATGTTGCCAAGGAA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       183 WWAWWAWWWATWWATCTATWTAAATAAATWTATAWWWAWAAAAAAWATAAAAA
                                                                                                                                                                                                                                                                                                                                                                                  348 TAAAGAAAACACTTAGATTCAATGATTGTAAATTTAAGGCAAATACACATATTAGTATT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       408 ACCTTAGTGTAATGTATCCCTGTCATATACAATAAGGTGAAATTATAAGTAC-CCTAT
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  527 ATCACTGGCTATGTGCTTAGATCTACAGGAGATCATATAATTTGATACAAATAAAAGAAA
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Drosophila melanogaster
Bukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
                                                                                                                                                                                                                                                                                                                              1;
                                                                                                                                                                                                                                                                Length 759;
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                                                                                                                                                                                                                                                                                                                          Mismatches 176;
organism="Yarrowia lipolytica"
                                                                                                                                                                                                                                                                   7.9%; Score 56; DB 9;
40.4%; Pred. No. 0.034;
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/clone="AW0AA009H09"
/clone_lib="AW0AA"
/note="end : T7"
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                           /mol_type="genomic DNA/strain="CLIB 89"
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487 (1), 52-55 (2000)
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Direct Submission
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Best Local Similarity
Matches 152; Conserv
                    20584718
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  isogenic strain y2; cn bw sp, the same strain used for the BDGP's Pl and EST libraries. A more detailed description of the library and how to order individual BAC clones, the entire library, or filters for hybridization from the BACPAC Resource Center can be found at http://bacpac.med.buffalo.edu/drosophila_bac.htm.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Souciet, J.L., Aigle, M., Artiguenave, F., Blandin, G., Bolotin-Fukuhara, M., Bon, E., Brottier, P., Casaregola, S., de-Montigny, J., Dujon, B., Durrens, P., Lepingle, A., Llorente, B., Malpertuy, A., Neuveglise, C., Ozier-Kalogeropoulos, O., Potier, S., Saurin, W., Tekala, F., Toffano-Nioche, C., Wesolowski-Louvel, M., Wincker, P. and Weissenbach, J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Zygosaccharomyces rouxii
Bukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
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                                                                                                                                                                                                                                                                                                                                                                   349 AAAGAAAAACACTTAGATTCAATGATTGTAAATTTAAGGCAAATACACATATTAGTATTA
                                                                                                                                                                                                                                                                                                                                                                                                     538 АААААЯКАТМАМАААТААТАААТАААААААМАТМААМИТМАНТММАТАТТТИМААААА
                                                                                                                                                                                                                                                                                                                                                                                                                                             409 CCTTAGTGTAATGTATCCCTGTCATATATACAATAAGGTGAAATTATAAGTACCCTATGC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   598 ИАМТТТСМААААТТМИААМТИМПААМТАИААТТИАТТАТИАУМАУААААААААА
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AGTIGGCIGGACAGTICTAAATIGGACTTTATTAATTTTTTAAAATCAGTAACTGATTTAT
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1 (bases 1 to 895)
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Wincker, P., Attiguenave, F. and Souciet, J.
Genomic exploration of the hemiascomycetous yeasts: 8.
Zygosaccharomyces rouxii
                                                                                                                                                                                                                                                                                       Length 996;
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                                                                                                                                                                                                                                                                                                                               45; Mismatches 141; Indels
                                                                                                                                  /organism="Drosophila melanogaster"
/mol_type="genomic DNA"
/db_xref="taxon:7227"
                                                                                                                                                                                                                                                                                         i; DB 9;
0.043;
                                                                                                                                                                                                                                                                                       7.8%; Score 55.6;
10.3%; Pred. No. 0.8
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                                                                                                                                                                                        /clone="BACR31021"
/clone_lib="RPCI-98"
/note="end : TET3"
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AUTHORS
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Submitted (NG-SEP-2000) Genoscope - Centre National de Sequencage, 2 rue Gaston Cremieux, CP 5706, 91057 EVRY cedex, FRANCE. (E-mail: sequefegenoscope.cns.fr - Web: www.genoscope.cns.fr)
This GSP is part of a random genomic sequencing program of thirteen yeast species: Saccharomyces bayanus var. uvarum, Saccharomyces excapanoyces bayanus var. uvarum, Saccharomyces cardaromyces servazzii, Zygosaccharomyces rouxii, Saccharomyces kluyveromyces thermocolerans, Kluyveromyces angusta, angusta, Barton Saccharomyces marxianus var. marxianus, Pichia angusta, Debaryomyces hansenii var. hansenii, Pichia sorbitophila, Candida tropicalis and Yarrowia lipolytica. Genomic inserts of 3 to 5 keywords for description of this sequence and for the sequence of the other extremity of this insert.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           510 TATATAATATYAATWATTAAAATAANAAAWWTTAAAATWATATATTATTAANTATAWATATA
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/evidence=not_experimental
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41.3%; Pred. No. 0.088;
tive 18; Mismatches 198; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        organism="Zygosaccharomyces rouxii"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               /mol_type="genomic DNA"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  'db_xref="taxon:4956"
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/product="tRNA-Thr"
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Submitted (12-APR-2000) Genoscope - Centre National de Sequencage
BP 191 91006 BVRY cedex - FRANCE (E-mail : seqref@genoscope.cns.fr
Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Actinopterygii, Neopterygii, Teleostei, Buteleostei, Neoteleostei,
Acanthomorpha, Acanthopterygii, Percomorpha, Tetraodontiformes,
Tetradontoidea, Tetraodontidae, Tetraodon.
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                                                                                                                                                                                                                                                                                                                                                                                compact genome of the
                                                                                                         Roest Crollius, H., Jaillon, O., Dasilva, C., Bouneau, L., Fisher, C., Bernot, A., Fizames, C., Wincker, P., Brottier, P., Quetier, F., Saurin, W. and Weissenbach, J.

Bstimate of human gene number provided by genome-wide analysis using Tetraodon nigroviridis DNA sequence
Nat. Genet. 25 (2), 235-238 (2000)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          556 AGATCATATAATTTGATACAAATAAAAGAAAAGTGTTCTCTCCCCTTACAGAATTGACAT
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                                                                                                                                                                                                                                                                                                         Roest Crollius, H., Jaillon, O., Dasilva, C., Ozouf-Costaz, C., Fizames, C., Fischer, C., Bouneau, L., Billault, A., Quetier, F., Saurin, W., Bernot, A. and Weissenbach, J. Characterization and repeat analysis of the compact genome of greshwater pufferfish Tetraodon nigroviridis
Genome Res. 10 (7), 939-949 (2000)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             /olone="082E19"
/clone_lib="G"
/note="Genoscope sequence ID : COBG082AC10SP1~end
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      This sequence is a single read and was generated as scale clone-end sequencing project of the Tetraodon genome. For more information, please take a look at http://www.genoscope.cns.fr/Tetraodon.
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44.0%; Pred. No. 0.14;
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/db_xref="taxon:99883"
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                                                                                                                                                                                                                                                                                         Web: www.genoscope.cns.fr)
Determination of this BAC-end sequence was carried out as part of a collaboration with the Buropean brosophila Genome Project (BDGP) - http://www.edgp.ebi.ac.uk -. This Drosophila melanogaster BAC library (Dros BAC) was made by Alain Billaud at CEPH (Centre d'Etdude du Polymorphisme Humain) with funding provided by a MRC project grant. The DNA was prepared from embryos by Alain Bucheton and Genevieve Payan. It has been constructed in the vector
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             787
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Submitted (23-JUL-1999) Genoscope - Centre National de Sequencage
BP 191 91006 EVRY cedex - FRANCE (E-mail : seqref@genoscope.cns.fr
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                846 WIWWATATATWIARRGGRAADDAAAAAATIWIIWIWWWWWWWIIWWWWWWWIIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            486 TAAATTGGACTTTAFTAATTTTTAAAATCAGTAACTGATTTATCACTGGCTATGTGCTTA
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Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
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/organism="Drosophila melanogaster"
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                                                                     Drosophila melanogaster (fruit fly)
Drosophila melanogaster
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37.2%; Pred. No. (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             /mol_type="genomic DNA"
/db_xref="taxon:7227"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         /clone="BACN11G11"
/clone lib="DrosBAC"
/plasmid="pBeloBAC11"
/note="end : T?"
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GSS; genome survey sequence.
Tetraodon nigroviridis
Tetraodon nigroviridis
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Best Local Similarity 37.2
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AL103735.1
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AC113817 Rattus no
AC126816 Rattus no
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AC094560 Rattus no
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AC150292 Callithri
                 AC024224 Homo sapi
AC006510 Homo sapi
AF400597 Homo sapi
E21012 Novel membr
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-Q=/cgn2_1/USPTO spool/US09989293/runat_01122004_102145_13797/app query.fasta_1.263
-DB=-GenEmbl - QPMT=fastap - SUFFTX=reg - MINMATCH=0. - LOOPELD=0 - LOOPEXT=0
-UNITS=bits -START=1 - END=-1 - MATRIX=blooum62 - TRANS=human40.cdi - LIST=1500
-UNITS=bits -START=1 - END=-1 - MATRIX=blooum62 - TRANS=human40.cdi - LIST=1500
-UNITS=bits -START=1 - END=-1 - MATRIX=D00 - THR MIN=0 - ALIGN=15 - NODE=LOCAL
-OUTFMT=pto -NORM=Ext - HEAPSIZE=500 - MINLEN=0 - MAXLEN=200000000
-USRR=US0989293 @CGN 1_1371 @runat_01122004_1021145_13797 - NOFU=6 - ICPU=3
-NO MMAP - LARGEQUERY - NEG SCORES - WAIT - DSPBELOCK=100 - LONGLOG
-PBV TIMEOUT=120 - WARN TIMEOUT=30 - THREADS=1 - XGAPOP=10 - XGAPEXT=0.5 - FGAPOP=6
-FGAPEXT=7 - YGAPOP=10 - YGAPEXT=0.5 - DELOP=6 - DELEXT=7
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AX055452 Sequence
AX403489 Sequence
AX464340 Sequence
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                             GenCore version 5.1.6
Copyright (c) 1993 - 2004 Compugen Ltd.
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66 14.3 2226307 10 ACI19851 66 14.3 2221682 2 ACI22603 66 14.3 222220 10 ACO91519 66 14.3 2310188 2 ACI31023 66 14.3 231018 2 ACI31023 66 14.3 231018 2 ACO91519 66 14.3 231010 2 ACO91519 66 14.3 233101 2 ACO91519 66 14.3 233101 2 ACO91519 66 14.3 233521 2 ACO91519 66 14.3 233521 2 ACO91519 66 14.3 233521 2 ACO91519 66 14.3 235521 2 ACO91519 66 14.3 235521 2 ACO91519 66 14.3 23621 2 ACO91519 66 14.3 23621 2 ACO9683 66 14.3 224360 2 ACI09684 66 14.3 224360 2 ACI09684 66 14.3 224360 2 ACI09684 66 14.3 22436 2 ACI09684 66 14.3 224467 2 ACI22764 66 14.3 224467 2 ACI22764 66 14.3 224467 2 ACI2268 66 14.3 224467 2 ACI2268 66 14.3 224467 2 ACI2268 66 14.3 22468 2 ACI36851 66 14.3 22468 2 ACI3685 66 14.3 22468 6 BDI10923 65.5 14.2 622 8 ACI3689 65.5 14.2 623 8 ACI3689 65.5 14.2 106 6 ACS5220	AC119851 Mus muscu AC12603 Rattus no AC11023 Mus muscu AC091519 Mus muscu AC091604 Mus muscu AC091604 Mus muscu AC13775 Rattus no AC095655 Rattus no AC126724 Rattus no AC126734 Rattus no AC126734 Rattus no AC126815 Rattus no AC094041 Rattus no AC094041 Rattus no AC02209 Homo sapi AC125831 Rattus no AC02209 Homo sapi AC109653 Rattus no AC096964 Rattus no AC109658 Rattus no AC109698 Rattus no AC109698 Rattus no AC109698 Rattus no AC109694 Rattus no AC102764 Rattus no AC122764 Rattus no AC122768 Rattus no AC122769 Rattus no	AC14931 Manan STS S AR415370 Sequence AX972240 Sequence BD110923 EST and e U67007 Stephanosph U66920 Aralia cali AF551720 Aralia cal BC07149 Homo sapi AX39625 Sequence AX06560 Sequence AX06661 Namnacara AX083665 Homo sapi AX03389 Biomphala AX03389 Biomphala AX03389 Biomphala AX03389 Biomphala AX03389 Biomphala AX03389 Biomphala AX0310126 Fusarium AX010809 Oryza sat AJ310126 Fusarium AX010809 Oryza sat AJ310126 Fusarium AR011338 Biomphala AX03389 Biomphala AX03389 Biomphala AX03389 Biomphala AX031389 Biomphala AX031389 Biomphala AX031389 Biomphala AX031389 Biomphala AX031389 Biomphala AX03128 Battus norv AB061437 Homo sapi S55323 gagrev (B06906 Visna virus AB001352 Leptospir AE001352 Leptospir AE0013537 S.pombe c BX649564 Mouse DNA AC017063 Homo sapi CONTINUALION (6 of
2 $4$ $4$ $4$ $4$ $4$ $4$ $4$ $4$ $4$ $4$	66 14.3 226307 10 66 14.3 228109 2 66 14.3 228109 2 66 14.3 229209 2 66 14.3 230193 2 66 14.3 230193 2 66 14.3 233141 2 66 14.3 233692 2 66 14.3 236921 2 66 14.3 22630 2 66 14.3 254360 2 66 14.3 271869 3 66 14.3 294467 2 66 14.3 29460 2 66 14.3 320054 2	66 14.3 324870 2 ACI40571 65.5 14.2 250 11 G15423 65.5 14.2 548 6 ARG1230 65.5 14.2 548 6 ARG7007 65.5 14.2 548 6 BD110923 65.5 14.2 600 8 SFU67006 65.5 14.2 602 8 ACI60520 65.5 14.2 623 8 ACI605920 65.5 14.2 623 8 ACI605920 65.5 14.2 10.8 9 BC007449 65.5 14.2 10.8 9 BC007449 65.5 14.2 10.8 9 BC007449 65.5 14.2 10.8 9 BC020749 65.5 14.2 10.8 9 BC020749 65.5 14.2 10.8 9 BC0207389 65.5 14.2 1137 5 AY030389 65.5 14.2 11314 3 AY0303389 65.5 14.2 1281 9 AY030399 65.5 14.2 1281 9 AY030399 65.5 14.2 2224 8 ACI01090 65.5 14.2 2224 8 ACI01090 65.5 14.2 3726 6 ACI01090 65.5 14.2 3726 10 ACI01090 65.5 14.2 11971 2 ACI01090 65.5 14.2 11971 2 ACI01090 65.5 14.2 11971 2 ACI01090 65.5 14.2 29676 8 SPACL486 65.5 14.2 29676 9 ACI010614-

RESULT 3  AX403489  LOCUS  DICCUS  DICCUS  AX403489  AX403489  AX403489. I G1:21436980  KENTANTON  AX403489. I G1:21436980  KENTANDS  SAME Home sapiens  GNGANISM  Mammalia, Butheria, Primates; Catarrhin; Hominidae; Home.  AUTHORS  ASHenara, A. Dasker, K.P., Botstein, D. Desnoyers, L., Eaton, D., Farrara, N., Caber, H., Gerriteen, M., Coddard, A., Codowski, P., Soni, NY.F., Roy, M., Sewart, T.A., Thumas, D., Watanabe, C.K., Hillams, P., Wood, W. I. and Zhang, T. A., Thumas, D., Watanabe, C.K., Hillams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Hillams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, P., Wood, W. I. and Zhang, T. Maranabe, C.K., Malinams, D., Watanabe, C.K., Malinams, P., Wood, W. I. and Shang, M. Maranabe, C.K., Malinams, M. Mollype="massigned blaw"   Additional Corner, Malinams, Malinams, M. Maranabe, C.K., Maranabe, C.K., Maranabe, C.K., Maranabe, C.K., Malinams, M. Maranabe, C.K., Maranabe, C.K., Malinams, M. Maranabe, C.K., Mara	
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 DEFINITION
 ACCESSION
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 gene
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

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Clark, H.F., Gurney, A.L., Abaya, E., Baker, K., Baldwin, D., Brush, J.,

Chen, J., Chow, B., Choic., Crowley, C., Currell, B., Deuel, B.,

Dowd, P., Eaton, D., Foster, J., Girmaldi, C., Gu, Q., Hass, P. E.,

Heldens, S., Huang, A., Kim, H.S., Klimowski, L., Jin, Y., Johnson, S.,

Lee, J., Lewis, L., Liao, D., Mark, M., Robbie, E., Sanchez, C.,

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Stinson, J., Vagts, A., Vandlen, R., Watanabe, C., Wieard, D., Woods, K.,

Xie, M.H., Yansura, D., Yi, S., Yu, G., Yuan, J., Zhang, M., Zhang, Z.,

Goddard, A., Wood, W.I. and Godowski, P.

The Secreted Protein Discovery Initiative (SPDI), a Large-Scale

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PUBMED
REFERENCE
AUTHORS
TITLE
JOURNAL
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AUTHORS
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
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B. (Dases 1 to 165414)

E. (Dases 1 to 165414)

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Muzny, D.M., Adams, C., Adio-Oduola, B., Ali-osman, F.R., Allen, C., Alsbrooks, S.L., Amaratunge, H.C., Are, J.R., Ayele, M., Banks, T., Barbaria, J., Benton, J., Bimage, K., Blankenburg, K. Bonnin, D., Borbaria, J., Benton, J., Bimage, K., Brown, M., Bryant, N.P., Bubay, C., Burch, P., Burkett, C., Brurell, K.L., Byrd, N.C., Carron, T.F., Carter, M., Cavazos, S.R., Chacko, J., Chavez, D., Chen, Z., Chiu, D., Chowdhry, T., Christopoulos, C., Cleveland, C.D., Cox, C., Coyle, M.D., Dathorne, S.R., David, R., David, R., David, M.L., David, M.L., David, M.L., David, K.R., Delgado, O., Denn, A.L., Ding, Y., Dinh, H.H.,
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Direct Submission Unpublished

(bases 1 to 165414) JOURNAL REFERENCE

Worley, K.C.

AUTHORS

Direct Submission

Submitted (28-FEB-2000) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA TITLE JOURNAL

3 (bases 1 to 165414) Worley, K.C.

AUTHORS JOURNAL

REFERENCE

Direct Submission

Submitted (26-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

(bases 1 to 165414)

Direct Submission Worley, K.C. REFERENCE AUTHORS

JOURNAL

COMMENT

Submitted (29-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 7030, USA
On May 29, 2002 this sequence version replaced gi:21206012.
INFORMATION: http://www.hgsc.bcm.tmc.edu/ or email gc-help@bcm.tmc.edu CLONE LENGTH: This sequence does not necessarily represent the entire insert of this clone. Overlapping regions of clones are only sequenced and submitted once, so the sequence for the remainder of the insert may be found in the record for the adjacent clones. Overlapping clones are noted at the beginning and end of the Features listing.

## ANNOTATION OF FEATURES:

STSs are identified using ePCR (Genome Res. 7:541-550) searches of a local database that includes entries from dbSTS, GDB, and local mapping efforts.

Repeats are identified using RepeatMasker (A. Smit and P. Green, unpublished.) for Human and Mouse sequences.
Genes and Region of sequence similarity are identified by BLAST (Nuc. Acids Res. 25:3389-3402) similarity (expect < 1e-34) to the EST and cDNA sequences. Genes demonstrate at least two exons flanked by consensus splice sites that maintained sequence

continuity across the splice junctions. Sequences that are not identical matches are annotated as similar.

SEQUENCING READ COVERAGE:Sequencing is completed to a minimum standard of double strand coverage with a minimum of 2 clones and 2 reads with no ambiguities or 2 chemistries with a minimum of 2 clones and 3 reads with no ambiguities. If the sequence quality for a region does not meet this standard, it will be indicated in the annotation as Low Coverage.

QUALITY OF INDIVIDUAL BASES: This sequence meets stringent quality standards - estimated error rate less than 1 per 10,000 bases. Reports of lowest quality individual bases and measures of base quality are listed below. Description of the metrics can be found

http://gc.bcm.tmc.edu:8088/quality.info/genbank.annotation.html.

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Malby, D.M., Adams, C., Adio-Oddola, B., Banks, T., Barks, T., Barks, J., Barks, T., Barks, J., Barks, T., Barks, J., Chavel, J., Chavel, J., Chavel, J., Chare, J., Chavel, J., Chare, J., Chavel, J., Dathorne, S.R., David, R., David, M. L., Davis, C., Coyle, M. D., Dathorne, S.R., David, R., David, M. L., Davis, C., Coyle, M. D., Dathorne, S.R., David, R., David, M. L., Davis, C., Earnhart, C., Edgar, D., Erdart, C., Edgar, D., Endard, C., Dugan, Rocha, S., Durbin, K.J., Earnhart, C., Edgar, D., Erdard, C., Escotto, M., Falls, T., Ferraguco, D., Flagg, N., Ford, J., Botter, P., Ferratz, P., Gabisi, A., Gao, J., Garcia, A., Ganzatne, P., Hamilton, K., Harris, C., Harris, K., Harris, M., Guraratne, P., Hanilton, K., Harris, C., Harris, K., Harris, M., Guraratne, P., Hanilton, K., Harris, C., Hadris, R., Hanilton, R., Jacobs, J., Jacobs, B., Jan, Y., Jooden, E., Moste, J., Jooden, S., Karlson, B., Jia, Y., Johnson, R., Jointes, M., Marinda, S., Karlson, B., Jia, Y., Johnson, R., Jointer, R., Hune, J., Louisegd, H., Lozado, R.J., Li, Z., Lichtarge, O., Lieu, C., Liu, J., Liu, M., Li, Z., Lichtarge, O., Lieu, C., Liu, J., Liu, M., Marinda, M., Mar
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
 Direct Submission
Submitted (05-FBB-1999) Molecular and Human Genetics, Baylor
College of Medicine, One Baylor Plaza, Houston, TX 77030, USF
On Sep 14, 2000 this sequence version replaced gi:10086352.
 Web site: http://www.hgsc.bcm.tmc.edu/
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Worley, K.C.
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JOURNAL
REFERENCE
AUTHORS
TITLE
ORGANISM
 JOURNAL
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; alternatively spliced"
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 Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JU-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
 1 (bases 1 to 625)
Willment, J.A., Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
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Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
 Villment, J.A., Gordon, S. and Brown, G.D.

Direct Submission
Submitted (18-JU-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
Location/Qualifiers
 1 (bases 1 to 744)
Willment,J.A., Gordon,S. and Brown,G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
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unclassified.

Es 1 (bases I to 741)

RS Shuji, H., Shoji, F., Kazunori, N. and Yasushi, A.

Shuji, H., Shoji, F., Kazunori, N. and Yasushi, A.

Novel membrane protein and DNA thereof

AL Patent: JP 1999001497-A 2 06-JAN-1999;

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PN JP 199901497-A/2

PD 06-JAN-1999

PF 31-JUN-1999

PF 31-JUN-1999

PF 31-JUN-1999

PF SHUJI HINUMA, SHOJI FUKUZUMI, KAZUNORI NISHI, YASUSHI ARAXA PC

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1. (bases 1 to 787)
Willment,J.A., Gordon,S. and Brown,G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms

J. Biol. Chem. 276 (47), 43818-43823 (2001)
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 20 AsnSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHis 39
 40 SerGinProThrGinSerSerLeuGluAspSerValThrProThrLysAlaValLysThr 59
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 Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUL-2001) Sir William Dunn School of Pathology,
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1 (bases I to 74)

Hermanz-Falcon, P., Arce, I., Roda-Navarro, P. and Fernandez-Ruiz, E. Cloning of human DECTIN-1, a novel C-type lectin-like receptor gene expressed on dendritic cells

Immunogenetics 53 (4), 288-295 (2001)
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 E 2 (bases 1 to 744)

S Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E.

B Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E.

Submitted (01-FBE-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain

B 3 (bases 1 to 744)

S Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E.

Direct Submission

Submitted (05-OCT-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain

Sequence update by submitter

On Oct 5, 2001 this sequence version replaced gi:14278818.
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87.10%
80.65%
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Best Local Similarity:
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 BD136999 1018 bp DNA linear PAT 18-SEP-2002 Isolated mammalian membrane protein gene and reagent relating
 G01N33/68,C12N15/00,A61K37/02,C12N5/00
Isolated mammalian membrane protein gene and reagent relating
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1 (bases 1 to 1018)
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Homo sapiens
Eukaryota, Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
 Sobanov, Y., Bernreiter, A., Derdak, S., Mechtcheriakova, D., Duechler, M., Kalthoff, F. and Hofer, E.
A novel cluster of lectin-like receptor genes expressed in monocytic, dendritic and endothelial cells maps close to the NK receptor genes in the human NK gene complex Unpublished
2 (bases 1 to 1153)
 de Martin,R., Hofer,E., Hofer-Warbinek,R., Kalthoff,F.S.,
Lipp,H.J., Mechtcheriakova,D., Schmid,J. and Sobanov,Y.
Ikk2 variant, dino gene, lectin-like receptor gene, and proteins
encoded thereby
Patent: WO 02077216-A 7 03-0CT-2002;
Novariis AG (CH); Novartis Erfindungen Verwaltungsgesellschaft
M.B.H. (AI)
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AUTHORS Sobanov,Y.

TITLE Direct Submitted (02-JUL-2011) Vascular Biology and Thrombosis Research,
JOURNAL Submitted (02-JUL-2011) Vascular Biology and Thrombosis Research,
University of Vienna, Brunnerstrasse 59, Vienna A-1235, Austria
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Alternative splicing: See also AJ312372.

Location/Qualifiers

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 Novel human secreted and transmembrane protein PRO1159 cDNA.
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US2003036179-A1.
 Human CDNA sequence encoding for PRO1159 polypeptide.WO200140466-A2.
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Mismatches:
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 Human PRO1159 (UN0589) nucleotide sequence SEQ WO200073454-Al.
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 cDNA encoding human PRO polypeptide #237.
 cDNA encoding human PRO1159 polypeptide.
US2002127576-Al.
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 ACD44377 standard; cDNA; 713 BP
 CDNA; 713 BP
 ACA03839 standard; cDNA; 713 BP
 Human secreted/transmembrane
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PN US200303531-A1.
PD 02-JAN-2003.
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Percent Similarity: 100.00%
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PA (GETH) GENENTECH INC.

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 DE Human secreted/transme PN US2003036179-A1. PD 20-FEB-2003. PA (GETH) GENENTECH INC. Percent Similarity: 100
 (GETH) GENENTECH INC.
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 07-DEC-2000.
(GETH) GENENTECH INC.
 ID ABX89377 standard; of Dr. DNA encoding novel s
PN US2003017563-A1.
PD 23-JAN-2003.
PA (GETH) GENENTECH IN
Percent Similarity: 1
Best Local Similarity: 1
 DNA encoding novel US2003017563-A1.
 ABX80868 standard;
 Percent Similarity:
Best Local Similarity:
 ACA64409 standard;
 Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
 AAF44240 standard;
 Percent Similarity:
Best Local Similarity:
 US2003036180-A1.
 US2003027162-A1.
 Percent Similarity:
 06-FEB-2003
 07-JUN-2001
 Query Match:
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RESULT 6
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 Query Match:
 RESULT 5
 GenCore version 5.1.6

Copyright (c) 1993 - 2004 Compugen Ltd.
- nucleic search, using frame plus p2n model

December 2, 2004, 05:01729; Search time 378 Seconds

1249.862 Million cell updates/sec
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Pred. No. 12: geneeqn2004s:*
score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
SUMMARIES
 462
1 MTFFLSLLLLLVCEAIWRSN.....DSRGLILGAEAWGRGVKKNT
 AACS6634 standard; cDNA; 713 BP.
Human PRO1159 protein UNQ589 encoding cDNA SEQ ID NO:272.
WO200053758-A2.
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 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Total number of hits satisfying chosen parameters:
 AAZ65094 standard; cDNA; 713 BP.
Membrane-bound protein PR01159 encoding cDNA
 Xgapop 10.0 , Xgapext 0.5
Ygapop 10.0 , Ygapext 0.5
Rgapop 6.0 , Fgapext 7.0
Delop 6.0 , Delext 7.0
4134886 seqs, 2624710521 residues
 Indels:
 Minimum DB seq length: 0
Maximum DB seq length: 200000000
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Maximum Match 100%
Listing first 1500 summaries
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 geneseqn2003cs:*
 AAC91481 standard; cDNA; 712 BP.
Human PRO1159 cDNA.
WO200073452-A2.
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nt Similarity: 100.00%

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 DE Human PRO1159 CDNA.
PN W0200073452-A2.
PD 07-DEC-2000.
PA (GETH) GENENTECH IN
Percent Similarity: 1
Best Local Similarity: 1
 Command line parameters:
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Best Local Similarity:
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 WO9963088-A2.
 14-SEP-2000
 09-DEC-1999
 Title:
Perfect score:
 Sequence:
Scoring table:
 Query Match:
RESULT 2
ID AAZ65094
DE Membrane
PN W0996308
PD 09-DEC-1
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Database :

Query Match: RESULT 3

RESULT 1

	PD 23-JAN Percent Sim Best Local Query Match	RESULT 21 ID ADA459 DE NOW459 PN US2003 PD 30-JAN	PA (GETH Percent Sim Best Local ( Query Match	RESULT 22 ID ADA7642 DE Human 1 PN US20030	PD 17-APR PA (GETH ) Percent Simi Best Local (	Query Match: RESULT 23 ID ADA1907 DE Human I	PN US2003( PD 20-MAR- PA (GETH ) Percent Simi	Best Local S Query Match: RESULT 24 ID HOAG165	PN US2003C PD 13-MAR- PA (GETH )	Percent Simi Best Local S Query Match: RESULT 25 ID ADB1948	DE Novel h PN US20030 PD 10-APR-	Siry Siry	· -	PA (GETH ) Percent Simi Best Local S Query Match: RESULT > ID ADA8650	DE Novel h PN US20030 PD 01-MAY-	rcer st 1 sty SUL7
Mismatches: 0 Indels: 0	otein cDNA, #154.	Conservative: 0 Mismatches: 0 Indels: 0	mbrane protein PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0	713 BP. transmembrane protein PRO1124 DNA.	Conservative: 0 Mismatches: 0 Indels: 0	ransmembrane protein, SBQ ID 473.	Conservative: 0 Mismatches: 0 Indels: 0	713 BP. transmembrane protein PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0		Conservative: 0 Mismatches: 0 Indels: 0	713 BP. transmembrane protein PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0	polypeptide PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0
Best Local Similarity: 100.00% Query Match: RESHE 19	ID ABX79548 standard; cDNA; 713 BP. DE Human sectreted/transmembrane protein PN 912202142961-A1.	3 t 2	ID ACA33569 standard; cDNA; 713 BP.  DB Novel human secreted and transmembrane protein PD 30-20202187-A1.	Percent Similarity: 100.00% Best Local Similarity: 100.00% Query Match: 100.00%	KESUL1 14 ID ARK81251 standard; DNA; 713 BP. DE Novel human secreted or transmem PN US20030427985-A1. PN OKE-PRE-2003	E T	1D ACAG4260 standard; cDNA; 713 BP. DE Human cDNA encoding a secreted/transmembrane by US203032155-A1.	iry iry	7 standard; cDNA; uman secreted and	ខ្លុំ ម៉ូ ម៉ូ		ង្គម្	6 standard; cDNA; uman secreted and	it it ge	standard; cDNA; 713 BP. creted and transmembrane 77615-A1.	PD 26-DEC-2002. PA (GETH ) GENENTECH INC. Percent Similarity: 100.00% Guery Match: 100.00% RESULT 20 ID ACJ81962 standard; CDNA; 713 BP.

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5992 standard; cDNA; 713 BP.

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14.) GENENTECH INC.

15. Anilarity: 100.00$ Mismatches: 0

16. Similarity: 00.00$ Indels: 0

17. Similarity: 00.00$ Conservative: 0

18. Similarity: 00.00$ Mismatches: 0

19. Similarity: 00.00$ Indels: 0
 181 standard; cDNA; 713 BP.
human secreted and transmembrane protein PRO1159 cDNA.
1068796-A1.
 01 standard, cDNA; 713 BP. human secreted and transmembrane protein PR01159 cDNA. 082711-A1.
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encoding human PRO1159 polypeptide.
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 122 standard; cDNA; 713 BP.
sncoding human PRO polypeptide #237.
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 073 standard; cDNA; 713 BP.
PRO polynucleotide #237.
3054517-A1.
 423 standard; cDNA; 713 BP. PRO polynucleotide #237. 3073212-A1.
 ADB16065 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003087350-A1.
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ರ.⊣	; cDNA; 713 BP. ng secreted/transmembrane	prot	ein PRO1159.	PA CGETH COUNTY OF PARCENT SIMILAR BEST LOCAL SIMILAR
8297-A1. 003. GENENTECH arity: milarity:	INC. 100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	Query Match: RESULT 38 ID ADA79465 E DE Human PRO PN US2030827
RESULT 30 ID ADA/381 standard; CDNA; DE Human PRO polynucleotide PN US20030/3212-A1.	cDNA; 713 BP.			it de
GENENTECH arity: milarity:	INC. 100.00\$ 100.00\$ 100.00\$	Conservative: Mismatches: Indels:	000	RESULT 39  ID ADA87604 8  ID ADA87604 8  ID NOVEL hum  PN US200330873  PD 08 AMY-200
ESULT 31  D ADA21573 standard; c  E Human cDNA encoding  N US2003054404-A1.	; cDNA; 713 BP. ng secreted/transmembrane		polypeptide PRO1159.	PA (GETH ) G Percent Similar Best Local Sim
PD 20-MAR-2003. Percent Similarity: Best Local Similarity: Query Match:	100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	ESSULT GEOMETRICATION OF THE PROPERTY OF THE P
RESULT 32 ID ADA10360 standard; of Human cDNA encoding PN US2003059831-A1.	; cDNA; 713 BP. ng secreted/transmembrane	prot	ein, PRO1159.	PN 022023087, PD 08-MAY-2087, PA (GETH ) G) Percent Simila.
ar ice	100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	Best Local Sim
RESULT 33 ID ADA67646 standard; DE Human PRO polynucle PN US2003068795-A1.	; cDNA; 713 BP. leotide #237.			PM US2003054 PD 20-MAR-20 PD 20-MAR-20 Percent Similar
03. ENENTECH rity: ilarity:	INC. 100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	 : 다음
RESULT 34  ID ADB30653 standard; c DE CDNA encoding human PN US2003068794-A1.	; cDNA; 713 BP. an PRO polypeptide #237	ide #237.		PN 02.002082 PD 01-MAY-20 PA (GETH ) G Percent Simila
10-APR-2003. (GETH ) GENENTECH coent Similarity: st Local Similarity: rry Match:	INC. 100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	Best Local Sim Query Match: RESULT 43 ID ADB1461 DE Human PRO
KESULI 33 ID ADA85949 standard; cDNA; 71 DE Novel human secreted and tr PN US2003082693-A1.	; cDNA; 713 BP. ted and transme	713 BP. transmembrane protein	PRO1159 CDNA.	PD 08-MAY-20 PA (GETH ) G Percent Simila
	INC. 100.00% 100.00% 100.00%	Conservative: Mismatches: Indels:	000	<del>1</del> ξ.Ε
KESULI 36 ID ADAI7904 standard; C DE cDNA encoding human PN US2003054987-A1.	; cDNA; 713 BP. an PRO1159 polypeptide	Peptide.		PD 17-APR-20 PD (7-APR-20 PA (GETH ) G Percent Simila Best Local Sim
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mman secreted and transmembrane protein PRO1159 cDNA.
13211-A1.
 standard; cDNA; 713 BP.
man secreted and transmembrane protein PRO1159 cDNA.
7345-A1.
 standard; cDNA; 713 BP.
man secreted and transmembrane protein PRO1159 cDNA.
12694-A1.
 standard; CDNA; 713 BP.

NA encoding secreted/transmembrane protein PRO1159.
4359-A1.
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O polynucleotide #237.
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 L standard; cDNA; 713 BP. NO polynucleotide #237. 37351-A1.
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ADB20033 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA. US2003082691-A1.
 ACD98660 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
 Human cDNA encoding secreted/transmembrane protein PR01159.
US2003059832-A1.
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 ADB24832 standard; cDNA; 713 BP.
Human PRO polynucleotide SEQ ID NO 473.
US2003077713-A1.
 Human PRO polynucleotide #237.
US2003082701-A1.
 ADA94137 standard; cDNA; 713 BP.
 ADB13345 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003082710-A1.
 ADA74599 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003068798-A1.
 ADA75319 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
 CDNA; 713 BP
 Human PRO polynucleotide #237.
US2003077722-A1.
DE Novel human secreteu con US2003082691-A1.
PD 01-MAY-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00%
Best Local Similarity: 100.00%
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PA (GETH) GENENTECH INC.
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 PD 10-APR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100
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(GETH) GENENTECH INC.
 24-APR-2003.
(GETH) GENENTECH INC.
 01-MAY-2003.
(GETH) GENENTECH INC.
 DE Human PRO polymuclee
PN US2003082701-A1.
PD 01-MAY-2003.
PA (GETH) GENENTECH IN
Percent Similarity: 1
Best Local Similarity: 1
 ADA94592 standard;
 Percent Similarity:
Best Local Similarity:
Query Match:
 Percent Similarity:
Best Local Similarity:
 27-MAR-2003
 Query Match:
RESULT 52
 Query Match:
RESULT 53
 RESULT 47
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Novel human secreted and transmembrane protein PRO1159 cDNA. US2003082695-A1.
 Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082708-A1.
 ADA38817 standard; cDNA; 713 BP.
Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003059780-A1.
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 ADB30101 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2003073214-A1.
 ADB25392 standard; cDNA; 713 BP.
Human PRO polynuclectide SEQ ID NO 473.
US2003077715-A1.
24-APR-2003.
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 CDNA; 713 BP
 CDNA; 713 BP.
 ADA84845 standard; cDNA; 713 BP
 Human PRO polynucleotide #237. US2005082703-A1.
 Human PRO polynucleotide #237.
US2003082761-A1.
 Human PRO polynucleotide #237. US200303210-A1.
 PN US200.50.-.
PD 17-APR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00%
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 PA (GETH) GENENTECH INC.
Percent Similarity: 100.00$
Query Match: 100.00$
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Percent Similarity: 100

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Percent Similarity: 100

Best Local Similarity: 100
 PN US2003082695-A1.
PD 01-MAY-2003.
PA (GETH) GENENTECH IN
Percent Similarity: 1
Best Local Similarity: 1
 ADA85397 standard;
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ADA75871 standard;
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ADA47096 standard;
 ADAB0629 standard;
 PD 27-MAR-2003.
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 01-MAY-2003
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Conservative: Mismatches: Indels:		Conservative: Mismatches: Indels:	ide #237.	Conservative: Mismatches: Indels:	ide #237.	Conservative: Mismatches: Indels:	nsmembrane protein	Conservative: Mismatches: Indels:		Conservative: Mismatches: Indels:	NO 473.	Conservative: Mismatches: Indels:	Conservative Mismatches: Indels:	Conservative:
INC. 100.00% 100.00% 100.00%	cDNA; 713 BP.	INC. 100.00% 100.00% 100.00%	cDNA; 713 BP. n PRO polypept:	INC. 100.00% 100.00% 100.00%	cDNA; 713 BP. n PRO polypept:	INC. 100.00\$ 100.00\$ 100.00\$	cDNA; 713 BP. g secreted/transmembrane	100.00% 100.00% 100.00%	CDNA; 713 BP.	INC. 100.00% 100.00% 100.00%	cDNA; 713 BP. eotide SEQ ID NO	INC. 100.00\$ 100.00\$ 100.00\$	cDNA; 713 BP. cotide #237. INC. 100.00% 100.00%	couna; 713 BP. leotide #237. INC.
<pre>BETH ) GENENTECH Similarity: cal Similarity: Agtch:</pre>	8 standard RO polynuci 77721-A1. 2003.	PA (GETH ) GENENTECH Dercent Similarity: Best Local Similarity: Query Match:	JULI 63 ADB26918 standard; cDNA; 713 BP. CDNA encoding human PRO polypeptide #23 VS200392147-A1.	<u> </u>	RESULT 84 RESULT	at Se	ID ADA92938 standard; cD DE Human cDNA encoding sPN US2003060407-A1.	PD 27-MAR-2003. Percent Similarity: Best Local Similarity: Query Match:	3 standard, piens. 49817-A1. 2003.		0 standard; RO polynucl 77714-A1.	24-APR-2003. (GETH) GENENTECH ccent Similarity: st Local Similarity: sty Match:	standard; O polynuch 22690-Al. 2003. GENENTECH arity: Imilarity:	ADAB1181 standard, ADAB1181 standard, Human FRC polynucl US2003082702-A1. 01-MAY-2003. (GETH ) GENENTECH rcent Similarity:
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Best Local Query Match RESULT 70 ID ADA99 DE Human PN US200 PN O1-MY PA (GETY PA (GE	PA (GE PACENT PERCENT BEST LOC QUERY MA RESULT 7 ID NOVE DE NOVE DE NOVE DE NOVE DE NOVE DE PACENT PACENT PACENT BEST LOC QUERY MA	ID AD DE HU DE HU DE PO 10 PA	PA (G) Percent Percent Best Lot Query M RESULT ID NO DE NO PN US PD 01 PA (G) PA (G) Percent Best Lot Query M	RESULT  1D AD  DE NO  DE NO  DE NO  DE NO  CO  DEST LO  DE NO  DE

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NA encoding secreted/transmembrane protein PRO1159.
9782-A1.
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creted/transmembrane PRO polypeptide cDNA #118.
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7712-A1.
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7982-A1.
 standard; cDNA; 713 BP.
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8793-A1.
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7711-A1.
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ADB46641 standard; cDNA; 713 BP.
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NDB35142 standard; cDNA; 713 BP.
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ADC57856 standard; cDNA; 713 BP.
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047 standard; cDNA; 713 BP.
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 92 standard; cDNA; 713 BP.
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 592 standard; cDNA; 713 BP.
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ID ADC56054 standard; cDNA; DE Novel human secreted and PN US20030937360-A1.	PA (GETHI 2003.  PA (GETHI 2003.  Percent Similarity:  Best Local Similarity:  Query Match:	KESULI 129 ID ADCS8624 standard; cDNA; E Novel human secreted and DN INSTANTALE_11	FN 08-MAY-2003. PD 08-MAY-2003. PA (GETH ) GENENTECH I Percent Similarity:	Best Local Similarity: Query Match:	KESULI 130 ADC14676 standard; cDNA; 713 BP. DE Novel human secreted and transmembrane protein PN US2003082546-Al.	ir ic	RESULT 131  ID ADD08208 standard; cDNA; DE Novel human secreted and PN US2003068623-A1.	PD 10-APR-2003. Percent Similarity: Best Local Similarity: Query Match:	NESULY 132 ID ADD03298 standard; cDNA; DE Novel human secreted and	2104-A1. 003. GENENTECH	Percent Similarity: Best Local Similarity: Query Match:	RESULT 133 ID ADC90290 standard; cDNA; DE Novel human secreted and	PN US20030B7348-Al. PD 08-MAY-2003. PA (GETH ) GENENTECH Percent Similarity:	ity:	ID ADC82033 standard; cDNA; DE Human PRO polymucleotide PN US2003084461-A1.	Percent Similarity: Best Local Similarity: Query Match:	RESULT 135 ID ADC69709 standard; cDNA; 713 BP. DE cDNA encoding human PRO polypeptide #237 PN US2003194770-Al.	16-CCT-Z003. (GETH) GENENTECH ccent Similarity: st Local Similarity: sty Match:	IE SOL 1898 Standard; CDNA; DE Human PRO polynucleotide PN US2003194773-Al.

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ADD55349 standard; cDNA; 713 BP. Human PRO polymucleotide #118. US2003077593-A1.		ESULT 154  ID ADD56307 standard; cDNA; 713 BP. DE Human PRO polynucleotide #118. PN US20030775594-Al.		11.155 MDD51875 standard; CDNA; 713 BP. CDNA encoding human PRO polypeptide UG200319479-A1.		ADD02614 standard; cDNA; 713 BP.  Human PRO polynucleotide #237.		ADD02108 standard; cDNA; 713 BP. Human PRO polynucleotide #237. 82508203430-Al.		I 158 ADD54290 standard; cDNA; 713 BP. Novel human secreted and transmembrane			NC. 100.00% 100.00%	cotide #237.  INC. 100.00\$ 100.00\$	A; 713 BP. de #237.
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ADE33518 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
VS2003194767-A1.
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ADESJA070 standard; cDNA; 713 BP.
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D ADEL9579 standard; CDNA; 713 BP.
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ADD93159 standard; CDNA; 713 BP.
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ABED9027 standard; cDNA; 713 BP.
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Best Local Similarity: Query Match:	RESULT 1/8 ADEZ2898 standard; CDNA; 713 BP. DE CDNA encoding human PRO polypeptide #237 PN VS2003199664-A1.	PA (GETH) GENERIECH Percent Similarity: Best Local Similarity: Query Match:	ID CDD79016 standard; DE CDNA encoding huma PN US2003203429-A1.	PD 30-OCT-2003. PA (GETH) GENENTECH INC. Percent Forcal Similarity: 100.00% Conservat Best Local Similarity: 100.00% Mismatche Query Match: 100.00% Indels:	KESULT 180 ID ADE26366 standard; c DE Novel human secreted PN US2003087305-A1.	PD 08-MAY-2003. Percent Similarity: Best Local Similarity: Query Match:	ID ADE32966 standard; DE Novel human secret	PD 16-OCT-2003. PA (GETH ) GENENTECH INC. Percent Similarity: 100.008 Best Local Similarity: 100.008 QUETY MAtch: 100.008 RESULT 182	ID ADE42658 standard; DB Human PRO polynucl. PN US2003199032-A1. PD 23-OCT-2003.	PA (GETH ) GENENTECH Percent Similarity: Best Local Similarity: Ouery Match: PERMIT 183	ID ADD80674 standard; cDNA; 713 BP. DE CDNA encoding human PRO polypeptide #? PN US2003207418-A1.	PD 06-NOV-2003. PA (GETH) GENEWIECH Percent Similarity: Best Local Similarity: Decy Local Similarity: Premm 184		ID ADE40986 standard; DE Human PRO polynucl PN US2003199031-A1. PD 23-OCT-2003. PA (GETH ) GENEWTECH Percent Similarity: Best Local Similarity: Query Match:

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D AD021523 standard; CDNA; 713 BP.

N Wovel human secreted and transmembrane protein PRO1159 CDNA.

N US2003207355-Al.

O 66-KNOV-2003.

A (GETH) GENEVIECH INC.

Sercent Similarity: 100.00$ Mismatches: 0

ESULT 190.
 PRO1159 cDNA.
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SULT 188
ADF67303 standard; CDNA; 713 BP.
Human PRO1159 nucleotide sequence SEQ ID NO:376.
 Conservative:
Mismatches:
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 A DG23264 standard; cDNA; 713 BP.

Novel human secreted and transmembrane protein
N US2003207384-Al.
O 06-NOV-2003.

A (GETH) GENENTECH INC.
STCent Similarity: 100.00$ Conservative: ser Local Similarity: 100.00$ Mismatches: lery Match: 101.00$
1D ADE04185 standard; cDNA; 713 BP.
DE Human PRO polynuclectide #237.
PN US2003199034-A1.
PD 23-CCT-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00$
Query Match:
RESULT 187.
 ID ADF97599 standard; cDNA; 713 BP.

DE Human PRO polynucleotide #237.

PN US200320730-A1.

PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00%

Query Match:

RESULT 192.
 PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00$

Query Match: 100.00$

RESULT 193

ID ADGB0111 standard; CDNA; 713 BP.

DE Human PRO polynuclectide #237.

PROSCOSSOSSOSSOSSOSSOSSOSSOSSOSSOSSOSSOSS
 ADE92914 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003194777-A1.
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 ADG80663 standard; cDNA; 713 BF
Human PRO polynucleotide #237.
US2003207373-A1.
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i (GETH) GENENTECH INC.

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(GETH) GENENTECH INC.

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PD 06-NOV-2003.  PA (GETH ) GENENTECH INC.  PACENT Similarity: 100.00\$ Conservative: 0  Percent Similarity: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Indels: 0  RESULT 203  ID ABX77952 standard; cDNA; 713 BP.  DE Human PRO polynclectide #118.  PN US203027163-A1.	st st sury sur	PN US20131222-A1. PN US201312222-A1. PD 19-SEP-2002. PA (GETH ) GENENTECH INC. Percent Similarity: 100.00\$ Mismatches: 0 Query Match: 100.00\$ Indels: 0 RESULT 205 ID ACA69270 standard; cDNA; 713 BP. DE Human CDNA nocoding secreted/transmembrane protein PRO1159.	PD 13-FEB-2003. PD 13-FEB-2003. Percent Similarity: 100.00\$ Conservative: 0 Best Local Similarity: 100.00\$ Mismatches: 0 Query Match: 100.00\$ Indels: 0 RESULT 206 ID ADD24089 standard; CDNA; 713 BP. DE Novel human secreted and transmembrane protein FR01159 CDNA.	요두권류	97,44	PN UG2002133125-A1.  PN UG202133125-A1.  PD 01-AUG-2002.  PA (GETH) GENENTECH LTD.  Percent Similarity: 100.00% Mismatches: 0  Best Local Similarity: 100.00% Mismatches: 0  Query Match: 100.00% Indels: 0  RESULT 209  ID ACA67230 standard; CDNA; 713 BP.  DE CDNA encoding human PRO polypeptide #237.  PN UG2003004311-A1.	PD 02-JAN-2003. PD 02-JAN-2003. PD (GETH) GENENTECH INC. PPATCAL Similarity: 100.00\$ Mismacches: 0 Beet Local Similarity: 100.00\$ Mismacches: 0 Query Match: 100.00\$ Indels: 0 RESULT 210 DE ADM82653 standard; CDNA; 713 BP. DE Novel human secreted and transmembrane protein PRO1159 CDNA. PN US2003087355-A1. PD 08-MAY-2003. PA (GETH) GENENTECH INC. Percent Similarity: 100.00\$ Conservative: 0
ADH55403 standard; cDNA; 713 Novel human secreted and tran US2003207381-A1. 06-NOV-2003. (GETH ) GENENTECH INC. rcent Similarity: 100.00\$ sty Local Similarity: 100.00\$ Sty Match: 195	13 BP. ransmemb	1: 100.00% 57 standard; CDNA; 713 BP. PRO polynuclectide #118. 050457-A1. 1-2003. illarity: 100.00% 11 arity: 100.00%	RESULT 197 ID AD164174 standard; CDNA; 713 BP. DE Novel human secreted and transmembrane protein PRO1159 CDNA. DE Novel human secreted and transmembrane protein PRO1159 CDNA. PN US2003207385-A1. PD 06-NOV-2003. PA (GETH ) GENENTECH INC. Percent Similarity: 100.00% Mismacches: 0 Query Match: 100.00% Indels: 0	standard; cDNA; 713 BP. uman secreted and transmemb 07386-A1. 2003. GENEWIECH INC. 100.00\$ imilarity: 100.00\$	100.00% Indels: d; cDNA; 713 BP. eted and transmembrane protein PR H INC. Conservative: 100.00% Mismatches:	100.00% Indels:  1; CDNA; 713 BP.  eted and transmembrane protein PR INC. 100.00% Mismacches: 100.00% Mismacches:	.3 BPansmembrane protein PRansmembrane protein PRmismatches: .Indels: .3 BPansmembrane protein PR.

Query Match: 100.00* Indels: 0 RESULT 219 ID ADE73974 standard; CDNA; 713 BP. DE Human PRO polynucleotide #237. PN US2003211571-A1.	PD 13-NOV-2003. PA (GETH ) GENBNTECH INC. Percent Similarity: 100.00% Conservative: 0 Best Local Similarity: 100.00% Mismatches: 0 Query Match: 100.00% Indels: 0 FOR MATCH CONSERVED CONS	DE CDNA encoding human PRO polypeptide #237.  PN US2003092108-A1.  PD 15-MAY-2003.  PA (GETH) GENENTECH INC.  Percent Similarity: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Indels: 0  RESULT 221.	ID ADE24002 standard; cDNA; 713 BP.  DE CDNA encoding human PRO polypeptide #237.  PN US2003092110-A1.  PD 15-MAY-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Mismatches: 0	ID ADE24645 standard; CDNA; 713 BP.  DE CDNA encoding human PRO polypeptide #237.  PN US2003092111-A1.  PD 15-MAY-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00% Mismatches: 0  Query Match: 100.00% Indels: 0  PERCENT 0.23	CDNA; 713 B ectide #237. INC. 100.00% 100.00%	CDNA; sotide INC. 100.00 100.00 100.00	# 237 # 237 # 237	US2003199054-Å1. 23-OCT-2003. (GETH ) GENENTECH INC. coent Similarity: 100.00% it Local Similarity: 100.00% iry Match:
<pre>% Mismatches: 0 /\$ Indels: 0 713 BP. transmembrane protein PRO1159 cDNA.</pre>	Conservative: 0 Mismatches: 0 Indels: 0	conservative: 0 Mismatches: 0 Indels: 0	13 BP.  transmembrane protein PRO1159 CDNA.  transmembrane protein PRO1159 CDNA.  Conservative: 0  mismatches: 0  Tndela: 0	embrane protein PR Conservative: Mismatches: Indels:	embrane protein PR Conservative: Mismatches: Indels:	Conservative: 0 Mismatches: 0 Indels: 0	Conservative: 0 Mismatches: 0 Indels: 0	Conservative: 0 Mismatches: 0
Best Local Similarity: 100.00% Query Match: RESULT 211 ID ADN16052 standard; CDNA; 713 BP DE Novel human secreted and transm PN US2003087351-A1.	08-MAY-2003. (GETH ) GENENTE (CCENT Similarity: St Local Similarity: ST MATCh: STUT 212	ID ADN16681 standard; CDNA; 713 BP.  DE Novel human secreted and transmembrane protein PP US203080395-A1.  PD 08-MAY-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00% Mismatches: Query Match: 100.00% Indels: Indels:	213 N15500 standard; CDNA; vvel human secreted and 22003087356-A1MAY-2003. ETH ) GENENTECH INC. Simlarity: 100.00 coal Similarity: 100.00 fatch: 100.00	948 standard; cDNA; human secreted and 3087357-A1. Y-2003. ) GENENTECH INC. milarity: 100.00 Similarity: 100.00		ADD76658 standard, cDNA, Human PRO polynucleotide US2003100087-A1. 29-MAY-2003. (GETH ) GENENTECH INC. rcent Similarity: 100.00 ery Match:	ADD80.22 standard; cDNA; Human PRO polymucleotide US200309213.A1. 15.MAY-2003. (GETH) GENENTECH INC. ccent Similarity: 100.00 sty Match: STLT 218 ADD86426_standard; cDNA;	DE Human PRO Polynucleotide #237. PN US-003203440-A1. PD 30-OCT-2003. PA (GETH) GENENTECH INC. Percent Similarity: 100.00%

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 ADE92362 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PR01159
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 CDNA encoding human PRO polypeptide #237.
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 cDNA encoding human PRO polypeptide #237.
US2003199029-A1.
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ADE94804 standard; CDNA; 713 BP.
CDNA encoding human PRO polypeptide #237.
US2003199027-A1.
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PD335502 standard; CDNA; 713 BP.
CDNA encoding human PRO1159 polypeptide.
US2003194760-A1.
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 ADE90663 standard; cDNA; 713 BP
 ADE91810 standard; cDNA; 713 BP
 CDNA; 713 BP.
 ADF35047 standard; cDNA; 713 BP
 ADE95356 standard; cDNA; 713 BP
 Human PRO polynucleotide #237.
US2003199063-A1.
 ADE93466 standard; cDNA; 713 B;
Human PRO polynucleotide #237.
US2003199060-A1.
 ID ADF35047 standalu,,
DE CDNA encoding human PRO polype
PN US203199029-A1.
PD 23-OCT-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00%
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Percent Similarity: 100
Best Local Similarity: 100
 DB ADF35502 standard; cDB cDNA encoding human PN US2003194760-A1.
PD 16-OCT-2003.
Percent Similarity: 1
Best Local Similarity: 1
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Best Local Similarity:
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 Query Match:
RESULT 235
ID ADE91810
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RESULT 234
ID ADE906
DE Human
PN US2003
PD 23-OCT
 Query Match
RESULT 230
RESULT 227
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DE Novel human secreted and transmembrane protein PR01159 cDNA.
PN US2003207360-A1.
PD 06-NOV-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00% Mismatches: 0
Best Local Similarity: 100.00% Mismatches: 0
DB Novel human secreted and transmembrane protein PRO1159 cDNA.
PN US2003199058-A1.
PD 23-007-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00% Mismatches: 0
Best Local Similarity: 100.00% Mismatches: 0
 ADG24368 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PR01159 cDNA.
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 RESULT 236
ID ADG11752 standard; CDNA; 713 BP.
DE CDNA encoding human PRO1159 polypeptide.
PN US2003228655.Al.
PD 11-DEC-2003.
PA (GETH) GENENTECH INC.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00% Mismatch
Best Local Similarity: 100.00% Mismatch
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 CDNA encoding human PRO polypeptide #237. US2003207376-A1.
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 ADG02389 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2002507352-A1. GENYV-2003. (GETH) GENENTECH INC.
 DE HUMAN PRO POLYMUZIOCIÓN, 1827.
PN 922003207422-A1.
PD 06-NOV-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00%
Best Local Similarity: 100.00%
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 ADF98151 standard; cDNA; 713 BP.
 ADG22175 standard; cDNA; 713 BP
 ADF98722 standard; cDNA; 713 Bl
Human PRO polynucleotide #237.
US2003208055-A1.
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RESULT 238
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 PA (GETH) GENENTECH INC.
Percent Similarity: 100
Best Local Similarity: 100
 06-NOV-2003.
(GETH) GENENTECH INC.
 Best Local Similarity:
 US2003207426-A1.
 ID ADG20245 standar
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NG2003207376-A1.
PD 06-NOV-2003.
PA (GETH) GENENTEC
Percent Similarity:
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 RESULT 242
 RESULT 243
 RESULT 237
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7351-A1. 003. GENENTECH : arity: milarity:	- P	RESULT 245  DE ADG16859 standard; CDNA; 713 BP.  DE CDNA encoding human PRO polypeptide #237  PN 022003207559-A1.  PN 06-NOV-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00% Conser  Best Local Similarity: 100.00% Mismat.  Query Match: 100.00% Indexp	RESULT 246  ID ADG05318 standard; CDNA; 7  DE Human PRO polynucleotide #  N U32003207375-A1.  PD 06-NOV-2003.  PA (GETH) GENENTECH INC.  Percent Similarity: 100.00%  Query Match: 100.00%	ADG19585 standard; ADG19585 standard; cDNA encoding humar US2003207425-A1.	PD 05-70V-2002 PA (GETH ) GENENTECH INC. Percent Similarity: 100.00\$ Conservat Best Local Similarity: 100.00\$ Mismatche Query Match: 100.00\$ Indels: RESULT 248	ADG13422 standard; CDNA encoding humar US2003207357-A1. 06-NOV-2003.	Percent Similarity:  Percent Similarity:  Best Local Similarity: 100.00%  Query March:  RESULT 249			ID ADG15649 standard; cDE cDNA encoding human PN US2003219885-A1.	PA (GETH) GENENTECH I Percent Similarity: Best Local Similarity: Ouery Match:	ADF97047 standard; cDNA; Human PRO polynucleotide US2003207371-A1.
PN PD PA Per Quest	DE D	PN P	RESI 1D DE PN PA Perc Perc	RESI ID DE PN	Perc Perc Best Ques	D D D D D	Per Per Best Que	DE PN	PD PA Perc Best Quer	PNE	Perc Perc Best Quer	S C B E

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Best Local Similarity: 100.00% Mismatches: 0

Best Local Similarity: 100.00% Mismatches: 0

Query Match: 100.00% Indels: 0

RESULT 256

ID MAG07303 standard; CDNA; 713 BP.

DE NOVel human secreted and transmembrane protein PRO1159 CDNA.

PN 052003207350-A1.

PN 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00% Mismatches: 0

Query Match: 100.00% Indels: 0

RESULT 257.
 PA (GETH) GENEVATOR.

PA (GETH) GENEVATION.

PA (GETH) GENEVATECH INC.

Best Local Similarity: 100.00% Mismatches: 0

Query Match: 100.00% Indels: 0

RESULT 258

ID ADG55350 standard; cDNA; 713 BP.

DE Novel human secreted and transmembrane protein PRO1159 cDNA.

PM US2003194778-A1.

PA (GETH) GENEVATION.

PA (GETH) GENEVATION.

Percent Similarity: 100.00% Mismatches: 0

Guery Match: 100.00% Indels: 0
 ADG23816 standard; cDNA; 713 BP. Novel human secreted and transmembrane protein PRO1159 cDNA. US2003207389-A1.
 ADG25006 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207427-A1.
 ADG07855 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002207356-A1.
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ADG61014 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
US200320/30-A1.
06-NOV-2003.
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 ADG06232 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003207374-A1.
 ADG04105 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003207423-A1.
 PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00%

Query Match: 100.00%

RESULT 253
PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00%

Best Local Similarity: 100.00%

Query Match: 100.00%
 PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00%

Query Match: 100.00%

RESULT 254
 PD 06-NOV-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 100.00%

Best Local Similarity: 100.00%

Query Match: 100.00%
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us-09-989-293a-377.rng.spdi

cent Similarity: 100.00\$  sty Match: 100.00\$  sty Match: 100.00\$  NUT 268  NOOE3 Standard; CDNA; 713  Novel human secreted and tran US2003207415-A1.  G-NOV-2003.  GETH ) GENENTECH INC.  CENT Similarity: 100.00\$  sty Match: 100.00\$	ID ADG71580 standard; CDNA; 713 BP.  DB Novel human secreted and transmembrane protein PRO1159 CDNA.  PN US2003207421-A1.  PD 06-NOV-2003.  PA (GETH ) GENEWIECH INC.  Percent Similarity: 100.00% Mismatches: 0  Best Local Similarity: 100.00% Mismatches: 0  RESULT 270  ID ADG81767 standard; CDNA; 713 BP.  DE Human PRO polynucleotide #237.	US2003207805-Å1.  06-NOV-2003.  cent Similarity: 100.00\$  it Local Similarity: 100.00\$  ityr 271  Human CDNA encoding secreted/ US200322865-A1.  1-DEC-2003.  (GETH ) GENENTECH INC.  cent Similarity: 100.00\$  ity Local Similarity: 100.00\$  ityr 272.	Standard, 20 Standard, 20 Standard, 27723-A1. 27723-A1. 27723-A1. GENEWIECH Larity: La	arity: 100.00% Conservative: milarity: 100.00% Mismatches: 100.00% Indels: standard; cDNA; 713 BP. man secreted and transmembrane protein PR 444.4A1. GENENTECH INC.	ccent Simil str Local Si str Local Si str Local Si SULT 275 ADG54246 Novel hu US200320 06-NOV-2 (GETH) >
PA (GETH) GENENTECH INC.  Percent Similarity: 100.00\$ Mismatches: 0  Guery Match: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Mismatches: 0  Guery Match: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Mismatches: 0	9 standard; cDNA; RO polynucleotide 07358-A1. GENENTECH INC. 1arity: 100.00 imilarity: 100.00	uman secreted and transmembrane p. 2003. 2003. 2003. GENENTECH INC. Larity: 100.00\$ Mismat. imilarity: 100.00\$ Mismat. Indels 6 standard; cDNA; 713 BP. Uman secreted and transmembrane p. 07364-A1. GENENTECH INC. CONSET. Larity: 100.00\$ Mismat. imilarity: 100.00\$ Mismat.	TOUR MEACH:  TOUR MACCH:  ADG55902 standard; CDNA; 713 BP.  ADG55902 standard; CDNA; 713 BP.  USO032037365-A1.  06-NOV-2003. (GETH) GENENTECH INC. (GETH) GENENTECH INC.  CORSETVATIVE:  TY MATCH:  TY MATCH:  TOUR MACCH:  TY MATCH:  TOUR MACCH:  TY MATCH:  TY MATCH:	PN 052003207358-A1.  PN 05-004-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00\$ Conservative: 0  Percent Similarity: 100.00\$ Mismatches: 0  Query Match: 100.00\$ Indels: 0  RESULT 266  ID ADG71028 standard; cDNA; 713 BP.  DE Novel human secreted and transmembrane protein PRO1159 cDNA.	dy t d

ID ADG59838 standard; cDNA; 71. DB Novel human secreted and tr	US2003207369-A1. 06-NOV-2003. (GETH) GENENTECH INC. cent Similarity: 100.00; st Local Similarity: 100.00; sry Match:	ਰਵੇਂ ਡ	cent Similarity: 100.00 tr Local Similarity: 100.00 try Match: 100.00 SULT 286 ADG10005 standard; CDNA;	DE Novel human secreted and try PP US2004009548-Al. PD 15-7AN-2004. PA (GETH ) GENENTECH INC. Percent Similarity: 100.00%	ery Match: SULT 287 ADI15476 standard	HINC.	terioral Similarity: 100.00 try Match: 100.00 sulf 288	LD AUGUSSSS REARGARG; CUNA; /LD DE NOVEL human secreted and tr. PN US2004009547-AL. PD 15-UNA-2004. PD 15-UNA-2004.	arity: 100.00 nilarity: 100.00	σō π	arity: nilarity: standard	eted and	arity: 100.00 nilarity: 100.00 100.00	1D ADJOSSES ESCRIBATIONS, 71.  DB NOVEL human secreted and try PD 26-FEB-2004.  PD 26-FEB-2004.  PACCENT ) GENENTECH INC.  Percent Similarity: 100.00% Query March: 100.00% RESULT 292  ID ADJ77579 standard; CDNA; 71.
Indels: 0	Conservative: 0 Mismatches: 0	is: protein PR	Conservative: 0 Mismatches: 0 Indels: 0	nne protein PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0	embrane protein PRO1159.	Conservative: 0 Mismatches: 0 Indels: 0	ne protein PRO1159 cDNA.	Conservative: 0 Mismatches: 0 Indels: 0	embrane protein PRO1159.	Conservative: 0 Mismatches: 0 Indels: 0		Conservative: 0 Mismatches: 0 Indels: 0	rane protein PRO1159 cDNA. Conservative: 0 Mismatches: 0
ery Match: 100.00% SULT 276	G81215 standard; CDNA; 713 BP. man PRO polynucleotide #237. 2003194793-A1CCT-2003. ETH ) GENENTECH INC. Similarity: 100.00\$	cd; cDNA; 713 BP.	06-NOV-2003. (GETH ) GENENTECH INC. rcent Similarity: 100.00% st Local Similarity: 100.00% PETY Match: 100.00%	0 standard, uman secret 07378-A1. 2003.	in ) Generater inc	KESOLI 4.7 BP.  ID ADH2115 standard; cDNA; 713 BP.  DE Human cDNA encoding secreted/transmembrane PN US2003224358-Al.	Similarity: 100.00% al Similarity: 100.00% tch: 100.00%	KESULT 280  DB AD561566 standard; CDNA; 713 BP.  DB Novel human secreted and transmembrane  PN US2003207429-A1.	PA (GETH) GENENTECH INC. Percent Similarity: 100.00% Cc Best Local Similarity: 100.00% Mi Query Match: 100.00% In	70.5	GERTH ) GENENTECH INC. GERTH ) GENENTECH INC. 100.00\$ st Local Similarity: 100.00\$ rry March: 100.00\$	က္ခြဲ	BETH ) GENENTECH INC. 2 similarity: 100.00% cal Similarity: 100.00% fatch: 100.00%	MESOLI 283  DB ADG54798 standard; CDNA; 713 BP.  DE Novel human secreted and transmembrane  BN US2003207367-A1.  PD 06-NOV-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 100.00% Cons.  Best Local Similarity: 100.00% Mism.  Ouery Match: 100.00% Indemism

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713 BP.
transmembrane protein PRO1159 cDNA.
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Mismatches:
Indels:
 713 BP.
olypeptide #237.
 113 BP.
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LEBECQUE S J E. PHILLIPS J H.
 CHALUS L.
QUAN A B.
BATES E E M.
GORMAN D M.
SAELAND S.
 Best Local Similarity:
PN US2003162955-A1.
PD 28-AUG-2003.
PA (CHAL/) CHALUS L
PA (QUAN/) QUAN A B
PA (BATE) BATES E
PA (GORM/) GORMAN
PA (GRM/) SAELAND
PA (LABE) LEBECQUE
PA (HALL/) PHILLIPS
PA (PHIL/) PHILLIPS
PECCENT SIMILATIVE
 DE LLR-J24-2 polyper
PN WO200277216-A2.
PD 03-CT-2002.
PA (NOVS) NOVARTIS
PA (NOVS) NOVARTIS
PA (NOVS) NOVARTIS
Percent Similarity:
 JS2003162955-A1.
 PD 03-OCT-2002.
PA (NOVS) NOVARTIS
PA (NOVS) NOVARTIS
Percent Similarity:
 Query Match:
RESULT 306
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RESULT 302
 Query Match:
RESULT 303
 Match:
 Query Matc
RESULT 301
 SEQ ID NO:36094
 ADF60937 standard; cDNA; 1018 BP.
Human long Schering dendritic cell membrane protein 4 (SDCMP4) cDNA.
 AAK81282 standard; DNA; 5709 BP.
Human immune/haematopoietic antigen genomic sequence
WO200157182-A2.
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 Conservative:
Mismatches:
Indels:
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 Mismatches:
 Indels:
 ADM27837 standard, cDNA, 713 BP.
cDNA encoding human PRO polypeptide #237.
US20040448333-A1.
 Indels:
 CDNA encoding human PRO polypeptide #237. US200403335-A1.
 cDNA encoding human PRO polypeptide #237.
US2004058424-A1.
25-MAR-2004.
 Indels:
 Indels:
 Indels:
 ADM28423 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2004077064-A1.
 DE Human SDCMP4 polypeptide encoding cDNA.
PN W09947673-A2.
PD 23-SEP-1999.
PA (SCHE) SCHERING CORP.
Percent Similarity: 87.10% Conse
Best Local Similarity: 80.65% Misma
 Jr 298
AAX01260 standard; cDNA; 741 BP.
Human DC3' protein coding sequence.
JP11001497-A.
 AAZ07533 standard; cDNA; 1018 BP.
 ADM42561 standard; cDNA; 713 BP
 DE CDNA enc.
DE CDNA enc.
DE CDNA enc.
PN US2004058424-AL.
PD 25-MAR-2004.
PA (GETH) GENEVITECH INC.
Percent Similarity: 100.00%

"T Local Similarity: 100.00%

"T Local Similarity: 100.00%
 Human PRO polynucleotide #237
 11-MAR-2004.
(GETH) GENENTECH INC.
(cent Similarity: 100.00$
st Local Similarity: 100.00$
 09-AUG-2001. (HUMA-) HUMAN GENOME SCI INC.
 PN OF-LAW-1999.
D 06-JAN-1999.
PA (TAKE) TAKEDA CHEM IND LTD.
Percent Similarity: 87.10%
Best Local Similarity: 80.65%
 Percent Similarity: 100.00%
Best Local Similarity: 100.00%
Query Match: 100.00%
 DE Human PRO polynucleotide #2
PN US200403836-A1.
PD 26-FEB-2004.
PA (GETH) GENENTECH INC.
Percent Similarity: 100.00%
Best Local Similarity: 100.00%
 Percent Similarity: 100.00%
Best Local Similarity: 100.00%
Query Match: 100.00%
 Percent Similarity: 100.00%
Best Local Similarity: 100.00%
Query Match: 100.00%
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 22-APR-2004.
(GETH) GENENTECH INC.
 26-FEB-2004.
(GETH) GENENTECH INC.
 Query March.
RESULT 293
Th ADJ65701 standard; of
 PN US2004077064-A1
PD 22-APR-2004.
PA (GETH) GENENTER
Percent Similarity:
 Query Match:
RESULT 297
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Human cDNA encoding a novel extracellular matrix protein, Seq ID No 223. WO200155368-A1.
 RESULT 307
ID AAK81284 standard, DNA, 336 BP.
DE Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36096.
 4 60 13
 000
 Conservative:
Mismatches:
Indels:
 MESULT 304

ID ADC10755 standard; cDNA; 1281 BP.

DE Human cDNA from extracellular matrix gene 66 #2.

PN US200359875-A1.

PD 27-MAR-2003.
Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
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Mismatches:
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 Mismatches:
Indels:
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 (NOVS) NOVARTIS AG. (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.
 (NOVS) NOVARTIS AG.
(NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.
 Indels:
 Indels:
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 Indels:
 RESULT 305
ID ABZ24041 standard; DNA; 1606 BP.
DE Human Dectin-1 polypeptide encoding DNA.
PN WO200296945-A2.
 LLR-J24-2 polypeptide encoding cDNA.
WO200277216-A2.
 ABV73363 standard; cDNA; 138 BP.
LLR-J24-stalk peptide encoding cDNA.
WO200277216-A2.
 Jr 303
ABG66733 standard; cDNA; 1281 BP.
Human polynucleotide SEQ ID NO 223.
US2002042386-A1.
 ABV73351 standard; cDNA; 1153 BP
 AAS31409 standard; cDNA; 1281 BP
 PN NCACALGE 2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 87.10%
Best Local Similarity: 80.65%
 DE Human polynucleotiue obc. I. S. 102002042386-Al.
PN 11-APR-2002.
PA (ROSE/) ROSEN C A.
PA (RARA/) BARAGH S M.
PA (BARA/) BARAGH S C.
PA (BARA/) BARAGH S C.
Percent Similarity: 87.10%
Best Local Similarity: 87.10%
 PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 87.10%
Best Local Similarity: 80.65%
Query Match: 53.57%
 PD 05-DEC-2002.
PA (ISIS-) ISIS INNOVATION LTD.
Percent Similarity: 87.10%
Best Local Similarity: 80.65%
 Percent Similarity: 100.00%
Best Local Similarity: 100.00%
Query Match: 50.65%
 80.65%
53.57%
 Percent Similarity: 87.10%
Best Local Similarity: 80.65%
Query Match: 53.57%
87.10%
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14 33 24

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Probe #12642 used to measure gene expression in human placenta sample.
WO200157272-A2.
09-AUG-2001.
 ID ABAJ1012 standard; DNA; 595 BP.

B Probe #9478 for gene expression analysis in human heart cell sample.

PN W0200157274-A2.

PD 09-AUG-2001.

PA (MOLE) MOLECULAR DYNAMICS INC.

Percent Similarity: 44.12% Mismatches: 33

Mismatches: 33
 Human genome-derived single exon probe from lung SEQ ID No 12059.
WO200186003-A2.
15-NOV-2001.
 AAK38061 standard; DNA; 595 BP.
Human bone marrow expressed single exon probe SEQ ID NO: 12618.
W0200157276-A2.
 Human foetal liver single exon nucleic acid probe #12136.
WO200157277-A2.
 AAK12342 standard; DNA; 595 BP.
Human brain expressed single exon probe SEQ ID NO: 12333.
WO200157275-A2.
 RESULT 323
ID AAS74556 standard; CDNA; 825 BP.
DE DNA encoding novel human diagnostic protein #10360.
 cDNA; 825 BP.
human diagnostic protein #24062.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
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Mismatches:
 Query Match:
RESULT 320
ID ABS37660 standard; DNA; 595 BP.
DE Human liver single exon probe, SEQ ID No 12670.
PN WO200157273-A2.
PN WO200157273-A2.
 Mismatches:
 Indel8:
 Indels:
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 PD 09-MUG-2001.

PA (MOLE-) MOLECULAR DYNAMICS INC.
Percent Similarity: 44.12*
Best Local Similarity: 30.39*
Query Match: 16.23*
 INC.
 INC.
 09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
 (MOLE-) MOLECULAR DYNAMICS INC.
 (MOLE-) MOLECULAR DYNAMICS INC.
 DNA; 595 BP
 DNA; 595 BP
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 43.66%
29.58%
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 PN WO200186003-A2.
PD 15-NOV-2001.
PA (MOLE-) MOLECULAR DYN Bercent Similarity: 44 Best Local Similarity: 36 Query Match:
RESULT 322
ID AASBR258 standard; CI DE DNA encoding novel hy WO200175067-A2.
 Percent Similarity:
Best Local Similarity:
3
Query Match:
1
RESULT 317
1D ABA31012 standard; D
E Probe #9478 for gene
PN WO200157274A2.
PD 09-AUG-2001.
PA (MOLE-) MOLECULAR DY
 Query Match:
RESULT 316
ID AA143956 standard; D
DE Probe #12642 used to
PN WO200157272-A2.
PD 09-AUG-2001.
PA (MOLE-) MOLECULAR DY
 ID AAS88258 standard, of DE DNA encoding novel he wo20017567-A2.

PD 11-OCT-2001.

PA (HYSE-) HYSEQ INC.
PRICERL Similarity: 4
Best Local Similarity: 2
Query Match:
 DE Human foetal liver in WO200157277-A2.
PD 09-AUG-2001.
PA (MOLE-) MOLECULAR DY Percent Similarity:
Best Local Similarity:
 RESULT 319
ID AAK12342 standard;
 ABS12068 standard;
 Percent Similarity:
Best Local Similarity:
Query Match:
 Percent Similarity:
Best Local Similarity:
 09-AUG-200
 Query Match:
 Query Match:
 analysis in human cervical cell sample.
 Human cDNA encoding a novel secreted protein, Seq ID 133. WO200155322-A2.
 111
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 AAC24137 standard; cDNA; 85 BP.
Human secreted protein 5' EST, SEQ ID NO: 28212.
EP10310-1A2.
06-58P-2000.
(GEST) GENSET.
 Conservative:
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 Mismatches:
 Mismatches:
Indels:
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 Indels:
 AAV42551 standard; cDNA; 528 BP.
Mouse dectin-1 extracellular domain cDNA.
WO9828332-A2.
 Mouse Dectin-1 polypeptide encoding DNA. W0200296945-A2.
 AAS25954 standard; cDNA; 1737 BP
 Probe #8766 for gene expression WO200157278-A2.
 ABZ24043 standard; DNA; 2298 BP
 ABX73295 standard; DNA; 1737 BP. Human novel polynucleotide #123.
US2002133753-A1.
19-SEP-2002.
 MOLEVALL 102-AUG-2001.

(HUMA.) HUMAN GENOME SCI INC. scent Similarity: 42.16% st Local Similarity: 30.39%
 (MOLE-) MOLECULAR DYNAMICS INC.
 AAI18833 standard; DNA; 595 BP.
 ABA63831 standard; DNA; 595 BP.
 ID ABZ24043 standard; DNA; 2298 BDE Mouse Dectin-1 polypeptide enc. PN W0200296945-A2.
PD 05-DEC-2002.
PA (ISIS-) ISIS INNOVATION LTD. Percent Similarity: 74.47% Best Local Similarity: 61.70% Query Match:
 AAV42548 standard; cDNA; 2298
Mouse dectin-1 cDNA.
WO9828332-A2.
 09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 ID AAV42548 standard; cDNA; 229
DE Mouse dectin-1 cDNA.
PN W0982832-A2.
PD 02-JUJ-1998.
PA (TEXA) UNIV TEXAS SYSTEM.
Percent Similarity: 74-47%
Best Local Similarity: 61.70%
Query Match: 31.06%
PN WC200157182-A2.
PD 09-AUG-2001.
PD (HUMA-) HUMAN GENOME SCI IN Percent Similarity: 100.00%
Best Local Similarity: 100.00%
 100.00%
 75.56%
62.22%
30.19%
 44.12%
30.39%
16.23%
 39.83%
 (TEXA) UNIV TEXAS SYSTEM
 26.41%
 42.16%
 Best Local Similarity: 30.39%
Query Match: 16.56%
 PN EP1033401-A2.
PD 06-SEP-2000.
PA (GEST) GENSET.
Percent Similarity: 1
Best Local Similarity: 1
 DE Human CDNA encoding
PN W0200155322-A2.
PD 02-AUG-2001.
PA (HUMA) HUMAN GENOME
Percent Similarity:
Best Local Similarity:
 PN W09828332-A2.
PD 02-JUL-1998.
PA (TEXA) UNIV TEXAS & Percent Similarity:
Best Local Similarity: (Query Match:
 PN WO200157278-A2.
PD 09-AUG-2001.
PA (MOLE-) MOLECULAR DY
Percent Similarity: 3
Best Local Similarity: 3
 (ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
 PN USZO02132753-AL
PD 19-SEP-2002
PA (ROSE/) ROSEN C
PA (RUBE/) RUBEN S
PA (BARA/) BARASH S
Percent Similarity:
 Query Match:
 Query Match:
RESULT 312
 Query Match:
RESULT 314
 Query Match:
RESULT 315
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14 33 24

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ABL18810 standard; DNA; 4367 BP.
 ID ABL18810 standar
DE Drosophila melan
PN W0200171042-A2.
PD 27-SEP-2001.
PA (PEKE) PE CORP
Percent Similarity:
 27-SEP-2001.
 Query Match:
RESULT 332
 Query Match:
RESULT 333
 Query Match:
 Query Match:
 RESULT 334
 Clone 040 ovule-specific gene encoding a cytochrome P450 monooxygenase.
US5907082-A.
 Query Match:
RESULT 330
ID ABL18811 standard; DNA; 2202 BP.
DD DE Drosophila melanogaster genomic polynucleotide SEQ ID NO 7906.
DP W0200171043-A2.
PD 27-SEP-2001.
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27
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
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Mismatches:
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 Conservative:
 ID ABWable
DB Human prostate expected
PN WO200160860-A2.
PD 23-AUG-2001.
PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Percent Similarity: 46.67% Conservative
Best Local Similarity: 29.33% Mismatches:
15.15% Indels:
 Mismatches:
Indels:
 Mismatches:
 Human prostate expression marker cDNA 49390.
WO200160860-A2.
 Indels:
 Indels:
 Indels:
 ADE84033 standard; DNA; 9289 BP.
5' regulatory region of human gene RBL2.
WO2003044226-A2.
 AAF11530 standard; cDNA; 629 BP.
Aspergillus niger EST SEQ ID NO:4053.
WO200056762-A2.
 Listeria innocua DNA sequence #1141.
 ABQ67682 standard; DNA; 771 BP.
Listeria innocua DNA sequence #484.
WO200228891-A2.
 DE Listeria innocua DNA sequence #484
DE Listeria innocua DNA sequence #484
PN WO20028891-A2.
PD 11-APR-2002.
PA (INSP) INST PASTEUR.
PA (CNRS) CNRS CENT NAT RECH SCI.
Percent Similarity: 31.37*
Best Local Similarity: 15.15*
 11-APR-2002.
(TRSP) INST PASTEUR.
(TRS) CONS CENT NAT RECH SCI.
cent Similarity: 47.06%
t Local Similarity: 31.37%
 (NOVO) NOVO NORDISK BIOTECH INC. (NOVO) NOVO NORDISK AS.
 AAX55600 standard; DNA; 1799 BP.
 T 326
ABV49399 standard; cDNA; 478 BP
 ABQ69702 standard; DNA; 771 BP
 ID ABL18811 standard; DNA; 220; DB Drosophila melanogaster gency WO20171042-A2.
PD 27-SED-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 50.79% Query Match: 15.15% RESULT 331
 PN 25-MAY-1999.
PD 25-MAY-1999.
PA (REGC) UNIV CALIFORNIA.
Percent Similarity: 43.21%
Best Local Similarity: 15.26%
 43.66%
29.58%
16.13%
 Percent Similarity: 48.28% Best Local Similarity: 29.31% Query Match: 15.37%
 Percent Similarity: 50.85%
Best Local Similarity: 27.12%
 30-MAY-2003.
(EPIG-) EPIGENOMICS AG.
PN WO200175067-A2.
PD 11-OCT-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity: 4
Best Local Similarity: 2
 WO200228891-A2.
 28-SEP-2000.
 11-APR-2002
 Query Match:
RESULT 329
 Match:
 Query Match:
RESULT 325
 Query Match:
RESULT 326
 Query Match:
RESULT 328
 Query Match
RESULT 324
 Percent
Best Loc
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Human cDNA #474 differentially expressed in activated vascular tissue. US2002137081-A1.
 AAF94482 standard; cDNA; 795 BP.
Human hydrophobic domain containing protein clone HP10715 cDNA #106.
22-FBB-2001.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 7903.
WO200171042-A2.
27-SEP-2001.
(PEKE) PE CORP NY.
 Drosophila melanogaster genomic polynucleotide SEQ ID NO 9721.
WO200171042-A2.
 11
21
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 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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 Mismatches:
 RESULT 335
ID ADLO8119 standard; DNA; 158091 BP.
DE Human gene associated with low HDL-C LIPC #1.
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 Listeria innocua contig DNA sequence #9.
WO200228891-A2.
 Rat gene AA893569, SEQ ID NO 6045. WO2003016475-A2.
 RESULT 336
ID ADF69677 standard; DNA; 181257 BP.
DE Human SLCSA8 gene SEQ ID NO:2.
PN WO2003104427-A2.
 PD 18-DEC-2003.
PA (UYCA-) UNIV CASE WESTERN RESERVE.
PA CHORD Similarity: 30.88*
 T 333
ABQ67196 standard; DNA; 684707 BP.
 ABL19416 standard; DNA; 4485 BP
 11-APR-2002.
(INSP) INST PASTEUR.
(CNRS) CNRS CENT NAT RECH SCI.
 CDNA; 771 BP
 PD 27-5E-5...

PA (PEKE) PE CORP NY.

Percent Similarity: 50 79%

Best Local Similarity: 31.75%

Best Local Similarity: 15.15%
 PA (VITI-) VIIIVALA 239.77*
Percent Similarity: 39.77*
Best Local Similarity: 27.27*
Best Local Similarity: 15.15*
 42.19%
34.38%
15.04%
 31.82%
15.04%
 Percent Similarity: 47.06$
Best Local Similarity: 31.37$
Onerv Match: 15.15$
 47.06%
31.37%
15.15%
 (GEHO) GEN HOSPITAL CORP. (FARB) BAYER AG.
 Percent Similarity: 50.79%
Best Local Similarity: 31.75%
Query Match: 15.15%
 15.15$
 53.03%
 04-MAR-2004.
(VITI-) VITIVITY INC.
 Percent Similarity:
Best Local Similarity:
Query Match:
 RESULT 337
ID ADE60145 standard;
 RESULT 338
ID ABX63474 standard;
 PD 26-SEP-2002.

PA (BAND/) BANDMAN O.

Percent Similarity:

Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
 US2004043389-A1.
```

Conservative: Mismatches: Indels:

us-09-989-293a-377.rng.spdi

(HELI-) HELIX RES INST. ccent Similarity: 53.03% 5t Local Similarity: 31.82% Bry Match: 15.04%	; cDNA; 2189 BP. cDNA clone SeqID BIOTECHNOLOGY.	arity: 53.03% milarity: 31.82% 15.04%		PA (PROT-) PROTEGENE INC.  Percent Similarity: 53.03\$ Conservati Best Local Similarity: 31.82\$ Mismatches Query Match: 15.04\$ Indels:	scandard, DNA, ecreted protein ge 63-A2.	Percent Similarity: 53.03% Mismatches:  Dest Local Similarity: 31.82% Mismatches:  Query March: 15.04% Indels:	ESOUL 331 ID ABB83426 standard; cDNA; 2271 BP. DE Human secreted protein gene 173 SEQ ID NO:24: PN WO200162891-A2.	PD 30-ANU-2001. PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 53.03* Conservati Best Local Similarity: 31.82* Mismatches Query Match: 15.04* Indels:	ESOUR 378 ACH04927 standard; cDNA; 2271 BP.  DE Novel human secreted protein #233 cDNA.  PN US2003065160-A1.  PD 03-APR-2003.	PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 53.03% Conservati: Best Local Similarity: 31.82% Mismatches Query Match: 15.04% Indels:	NA; L sec	Percent Similarity: 53.03\$  Best Local Similarity: 31.82\$  Mismatches: 15.04\$  RESULT: 354  RESULT: 354	ID AAV04583 standard; DNA; 2276 BP.  BE Human secreted protein gene 173 clone HEBMG65 PN WO9854963-A2. PD 10-DEC-1998.  PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 53.03
Conservative: 14 Mismatches: 22 Indels: 9	sequence, SEQ ID NO: 1972.	Conservative: 14 Mismatches: 22 Indels: 9	ID NO: 475.	Conservative: 14 Mismatches: 22 Indels: 9	SeqID 475.	Conservative: 14 Mismatches: 22 Indels: 9	CDNA cluster SeqID 1972.	Conservative: 14 Mismatches: 22 Indels: 9		Conservative: 14 Mismatches: 22 Indels: 9	tic protein #17357.	Conservative: 11 Mismatches: 16 Indels: 5	P.  HUMAN GENE GROUP.  CONSERVATIVE: 14  Mismatches: 22  Indels: 9  IP.  NO: 3071.
AGA ) SAGAMI CHEM RES CENT. ROT-) PROTEGENE INC. 53.03* cal Similarity: 53.03* latch:	2 standard; cDNA; 923 BP. DNA clone representative 94.A2. 2001.	(HELL-) HELLX RES INST. cont Similarity: 53.03% tt Local Similarity: 31.82% sry Match: 15.04%	SULT 341 AAK92015 standard; cDNA; 923 BP. Human cDNA 5'-end sequence, SEQ EP1130094-A2. 05-SEP-2001.	(HELL-) HELLX RES INST. ccent Similarity: 53.03\$ st Local Similarity: 31.82\$ sry March: 15.04\$ sulf 342	Ablicates scandard; cons, 523 br. 5' end of a human cDNA molecule EP1396543-A2.	Central : Assacramentation: Central : 53.03% it Local Similarity: 53.03% it Local Similarity: 31.82% its Match: 15.04%	ard; cDNA; 923 BP. presentative human	PD 10-MAK-2004. PA (REAS-) RES ASSOC BIOTECHNOLOGY. Percent Similarity: 53.03* Best Local Similarity: 31.82* Query Match: 15.04*	ID ABQ9935 standard, cDNA, 971 BP. DE Human coding sequence SEQ ID 58. PN WC200259566-A2. PD 01-AUG-2002.	(HYSE-) HYSEQ INC. cent Similarity: 53.03% it Local Similarity: 31.82% sry Match: 15.04%	cDNA; 1045 BP human diagnos	arity: 57.14% milarity: 34.69% 15.04%	ID AAF74861 standard; cDNA; 1999 BP. DE Human hTGN48-1so mucleotide sequence PN CN1272541-A. PD 08-NGV-2000. PA (NANF-) NANFANG RES CENT STATE HUMAN RECORT SImilarity: 33.03% Query Match: 15.04% RESULT 347 ID AAK94357 standard; cDNA; 2189 BP. DE Human full-length cDNA; 2189 BP. PN EP1130094-A2. PD 05-SEP-2001.

OLOGY. Conservative: 14 Mismatches: 22 Indels: 9	2198 BP. containing protein clone HP10715 cDNA #116.	CENT.  Conservative: 14  Mismatches: 22  Indels: 9	2271 BP.  gene 173 clone HE8MG65.  I INC.  Conservative: 14  Mismatches: 22 144* Indels: 9	2271 BP. ne 173 SEQ ID NO:243. INC. Conservative: 14 Mismatches: 22 Indels: 9	2271 BP. ein #233 cDNA. INC. Conservative: 14 Mismatches: 22 Indels: 9	A; 2271 BP.  secreted protein gene 173 #2.  CI INC.  CONSERVATIVE: 14  Mismatches: 22  82\$  Tindels: 9	76 BP.  10 173 clone HE8MG65.  10 Conservative: Mismatches: Indels:
PD 10-MAR-2004. PA (REAS-) RES ASSOC BIOTECHNOLOGY Percent Similarity: 53.03% Best Local Similarity: 31.82% Query Match: 15.04%	2 standard; ydrophobic d 2660-A2.	(SAGA ) SAGAMI CHEM RES (SAGA ) SAGAMI CHEM RES (PROT-) PROTEGENE INC. Froeth Similarity: 53.02 St Local Similarity: 31.82 ST MATCH: 15.04	2 standard; DNA; ecreted protein g 63-A2. 1998. HUMAN GENOME SCI larity: 31 82. imilarity: 15.04	RESULT 351 ID ABAB3426 standard; CDNA; 22 DE Human secreted protein gene PN W0200162891-A2. PD 30-AUG-2001. PA (HUMA-) HUMAN GENOME SCI IN Percent Similarity: 53.03% Best Local Similarity: 31.82% Ouery March: 15.04%	ESCUL 327.  DE ACH04927 standard; CDNA, 227.  DE Novel human secreted protein  PN US2003065160-A1.  PD 03-APR-2003.  PA (HUMA-) HUMAN GENOME SCI INC  Percent Simlarity: 53.03%  Best Local Similarity: 15.04%  OMERY Match:	7 standard; cDNA from novel 74-B1. 2003. HUMAN GENOME S larity: 53. imilarity: 15	standard; DNA; ecreted protein g 63-A2. 1998. HUMAN GENOME SCI larity: 53.03 imilarity: 31.82

Indels:

```
DE Drosophila melancy expressed polynucleotide SEQ ID NO 15770.

PN W020071042-A2.

PD 27-SEP-2001.

PA (PEKE) PE CORP NY.

Percent Similarity: 45.56$ Conservative: 15

Best Local Similarity: 28.89$ Mismatches: 24

RESULT 371
 AAZ93151 standard; DNA; 1921 BP.
Sequence encoding DNA replication and repair associated protein.
WO200008156-A2.
 SEQ ID NO 37579.
 13
25
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25
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23
17
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23
17
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 ABL28702 standard; DNA; 3813 BP.
Drosophila melanogaster genomic polynucleotide
WO200171042-A2.
 DE Toxicity modelling related rat gene SEQ ID No PN W020025000-A2.
PD 28-NOV-2002.
PA (GENE-) GENE LOGIC INC.
Percent Similarity: 47.37% Conservative Best Local Similarity: 27.63% Mismatches: RESULT 370
 Mismatches:
Indels:
 Mismatches:
Indels:
 RESULT 368

ID ABG22564 standard; CDNA; 3413 BP.

DE RAT CD71 encoding CDNA SEQ ID NO:27.

PN WO2002100336-A2.

PD 19-DEC-2002.

PA (TARG-) TARGET PROTEIN TECHNOLOGIES INC. Percent Similarity: 47.37% Conservat Best Local Similarity: 27.63% Mismatche Query Match: 369
 ID AAH1G060 standard; cDNA; 2539 BP.

DE Human cDNA sequence SEQ ID NO:14742.

PN EP1074617-A2.

PD 07-FEB-2001.

PA (HELI-) HELLX RES INST.

Percent Similarity: 51.61% Conse Best Local Similarity: 30.65% Misma Query Match: 14.83% Indell RESULT 367

ID AAH18122 standard; cDNA; 2574 BP.

DE Human cDNA sequence SEQ ID NO:17987.
 Query Match: 14.83* In
RESULT 366
ID AAH16060 standard; CDNA; 2539 BP.
DE Human CDNA sequence SEQ ID NO:14742.
PN EP1074617-A2.
 RESULT 364

ID ABX73737 standard; DNA; 1742 BP.

DE Human novel polymucleotide #565.

PN US2002132753-A1.

PD 19-SEP-2002.

PA (ROSE/) ROSEN C A.

PA (RUBE/) RUBEN S M.

PA (BARA/) BARASH S C.

Percent Similarity: 43.27%

Best Local Similarity: 30.77%
 ABL07096 standard; cDNA; 3810 BP
 D AAZ93151 standard; DNA, LOLD B Sequence encoding DNA replica PN WO20000156-A2.

PD 17-FEB-2000.

PA (INCY-) INCYTE PHARM INC. Percent Similarity: 51.61% Best Local Similarity: 30.65%
 ID AAH18122 standard; CDNA; 257
DE Human CDNA sequence SEQ ID N
PN EP107461-A2.
PD 07-FEB-2001.
PA (HELL:) HELLX RES INST.
Percent Similarity: 51.61%
Best Local Similarity: 30.65%
Query Match: 14.83%
14.83%
 45.56%
 PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity:
 Query Match:
RESULT 365
Query Match:
```

	Mismatches: 24 Indels: 25		_
5 BP. omic p	5875 BP. genomic polynucleotide SEQ ID NO	NO 27571.	
	Conservative: 24 Mismatches: 33 Indels: 17		
DNA; 6240 BP. Ister genomic p	6240 BP. genomic polynucleotide SEQ ID	) NO 3649.	
	Conservative: 15 Mismatches: 24 Indels: 25		
3 BP. #35021.			
	Conservative: 17 Mismatches: 20 Indels: 22		
1206 BP. diagnost	1206 BP. diagnostic protein #28314.		
	Conservative: 12 Mismatches: 31 Indels: 1		
cDNA; 1681 BP.			
SHANGHAI Con Mis	HAI. Conservative: 12 Mismatches: 32 Indels: 30		
cDNA; 1990 BP. 3 clone PLACE60	:DNA; 1990 BP. clone PLACE60175640.		
INST. BIOTECHNOLOGY. 44.30% 25.32% 14.72%	Conservative: 15 Mismatches: 34 Indels: 10		
ADE07768 standard; DNA; 2346 BP. Novel coding sequence (useful for MO2003054122-A2.	ADE/7768 standard; DNA; 2346 BP. Novel coding sequence (useful for identifying genetic MO2003054152-A2.	disorders) #834.	
	Conservative: 11 Mismatches: 28 Indels: 26		
2625 BP. BGL3 enc	BP. encoding sequence.		
	Conservative: 20		

sequence.	d cells SeqID
α ου ο ωνο νου · ασυ	vate vate 79.
smatches: ength encodi smatches: smatches: dels: smatches: dels: smatches: dels: smatches: smatches: smatches: dels: smatches: dels: smatches: dels:	INE INC.  INE INC.  ISMATCHES:  IGHIS:  IN MYCN ACT:  ISMATCHES:  IGHIS:  INC. 299
TY: 25.97* Mi  ard; CDNA; 2785 BP. Ir  2.  OR INT INC. CC  EY: 25.97* Mi  14.72* Ir  ard; DNA; 5282 BP. Ir  ard; DNA; 5282 BP. Ir  GENOME SCI INC. CC  ty: 23.97* Ir  ard; DNA; 849 BP. CC  ty: 23.97* Mi  ard; DNA; 849 BP. CC  ty: 24.49* Mi  ard; CDNA; 1326 BP. CC  ty: 24.59* Mi  ard; CDNA; 1388 BP. CC  ty: 27.59* Mi  ard; CDNA; 1398 BP. Ir  ard; CDNA; 1398 BP. CC  ty: 27.59* Mi	mark 111VE expr exps exps exps exps 1699 NA #6
Best Local Similarity: 25.97% Query Match:  RESULT 380  DE ACCS0041 standard; CDNA; 278; DE Novel beta-galactosidase BGL; DN 03-APR-2003.  PA GEMV) GENBUCK INT INC. Percent Similarity: 25.97% Query Match: 14.72% RESULT 381  D AAK80242 standard; DNA; 5282 DN WO200157182-A2. PA (HUMA-) HUMAN GENOME SCI INC Percent Similarity: 23.97% Query Match: 14.72% RESULT 381  DD 09-AUG-2001. PA (HUMA-) HUMAN GENOME SCI INC Percent Similarity: 23.97% Query Match: 14.72% RESULT 382  DD 8-AUG-2001.  DE E faccium DNA; 849; DD 24-JUN-2003. PN (SE68275-B1) PN 40568275-B1 PN 40568275-B1 DD AAS78554 standard; CDNA; 1320 DE DNA encoding novel human diagent MO200175067-A2. DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA encoding novel human diagent MACON-15081 (14.61% RESULT 383 DE DNA ENCON-15081 (14.61% RESULT MACON-15081 (14	Human prostate expy WO200160860-A2. 23-AUG-2001. (MILL) MILLENNIUM reent Similarity: st Local Similarity: st Local Similarity: US2003119009-A1. SULT 385 (NUCH) NUCHTERN J (NUCH) NUCHTERN J (STUA/) STUART S G (STUA) STUART S G (STUART S G (STU

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Query Match:
RESULT 396
 Query Match:
RESULT 399
 Query Match:
RESULT 402
 RESULT 397
 RESULT 398
 Human secreted protein clone bf157_16 nucleotide sequence SEQ ID NO:93. WO200009552-A1.
 Base sequence of DNA fragment containing yeast URA3 gene.
WO9807873-A1.
26-FEB-1998.
 13
31
27
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31
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31
27
 14
42
8
 10
42
28
 14
42
 Arabidopsis thallana DNA; 1418 BP.
BP103345-A2.
G6-SEP-2000.
 cDNA; 2430 BP.
human diagnostic protein #10598.
 cDNA; 2991 BP. human diagnostic protein #27796.
 cDNA; 2991 BP. human diagnostic protein #9464.
 Conservative:
Mismatches:
Indels:
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 Indels:
 Human polynucleotide SEQ ID NO 325.
W0200153312-A1.
26-JUL-2001.
GHYSE-) HYSEQ INC.
 AAV31505 standard; DNA; 2330 BP.
 DNA; 2330 BP
 AAA16664 standard; cDNA; 2922
 PD 11-007-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity: 26.44%
Best Local Similarity: 26.44%
 PA (H1502) 43.69%
Percent Similarity: 43.69%
Best Local Similarity: 31.07%
 43.69%
31.07%
14.61%
 36.94%
27.93%
14.50%
 ent Similarity: 43.69%
Local Similarity: 31.07%
Match: 14.61%
 Paragram (GEMY) GENETICS INST INC.
Percent Similarity: 43.69%
Best Local Similarity: 31.07%
Ouery Match: 14.61%
 26.44%
14.61%
 42.53%
 ID
DE Hum.
PN WG2015.
PD 26-JUL-2001.
PA (HYSE-) HYSEQ 1.
Percent Similarity:
"atch: standard 1 nov
 PETERSON D P.
 COCKS B G.
HAWKINS P R.
 LT 391
AAS91992 standard; c
DNA encoding novel h
WO200175067-A2.
 AAS74794 standard;
DNA encoding novel
WO200175067-A2.
 Query Match:
RESULT 393
ID AAC41257 standard;
 PD 11-OCT-2001.

PA (HYSE-) HYSEQ INC.

Percent Similarity:

Best Local Similarity:
 AAS73660 standard;
 DNA encoding novel WO200175067-A2.
 AAT08603 standard;
 11-OCT-2001.
(HYSE-) HYSEQ INC.
 Best Local Similarity:
PA (PETE/) PETERSON
PA (COCK/) COCKS B
PA (HAWK/) HAWKINS
PETCENT SIMILATILY;
BEST LOCAL SIMILATILY;
 Percent Similarity:
 24-FEB-2000
 Query Match:
RESULT 391
 Query Match:
RESULT 392
 Query Match:
RESULT 394
 Query Match
RESULT 390
 Query M
RESULT
 Query M
RESULT
ID AA
DE BA
PN WO
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Oligonucleotide for detecting cytosine methylation SEQ ID NO 39839. WO200218632-A2.
 ABQ53249 standard; DNA; 724 BP. Oligonucleotide for detecting cytosine methylation SEQ ID NO 39840.
WO200218632-A2.
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:45. WO2003048304-A2.
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:47.
W02003048304-A2.
12-UIN-2003.
(AMHP) WYETH HOLDINGS CORP.
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 Conservative:
Mismatches:
Indels:
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Mismatches:
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 Mismatches:
 Mismatches:
Indels:
 Mismatches:
Indels:
 Mismatches:
Indels:
 Pseudomonas aeruginosa polynucleotide #16320.
US6551795-B1.
 ABD17357 standard; DNA; 1332 BP.
Pseudomonas aeruginosa polynucleotide #15961.
 Indels:
 Indels:
 Indels:
 Indels:
 PD 03-JAN-2003.

PA (SYGN) SYNCENTA PARTICIPATIONS AG.

Percent Similarity: 48.44% C.

Best Local Similarity: 40.62% M

Query Match: 14.39% II
 PN UNDEADLY.

PD 22-APR-2003.

PA (GENO-) GENOME THERAPEUTICS CORP.

Percent Similarity: 41.89%

Best Local Similarity: 25.68%
 PA (GENO.) GENOME THERAPEUTICS CORP.
Percent Similarity: 41.89%
Best Local Similarity: 25.68%
Query Match: 14.39%
RESULT 400
 DNA; 3636 BP.
SEQ ID NO 7775.
 ADB06105 standard; DNA; 1611 BP
 BP
 ADA72740 standard; DNA; 2000 BP.
Rice gene, SEQ ID 6065.
WO2003000898-A1.
 ABD17716 standard; DNA; 1065 BP
 DNA; 724 BP
 Ouery Match: 14.39%
RESULT 401
ID ADB06107 standard; DNA; 1611
 ID ABQ53249 standaru,,
DE Oligonuclectide for detecting
PN W0200218632-A2.
PD 07-MAR-2002.
PA (EPIG-) EPIGENOMICS AG.
Percent Similarity: 47.92%
Best Local Similarity: 43.75%
 PD 12-JUN-2003.

PP (APPL) WYETH HOLDINGS CORP.
Percent Similarity: 43.06%
Best Local Similarity: 30.56%
 RESULT 403

ID ADE61846 standard; DNA; 3636

DE Human gene M22919, SEQ ID N
PN WC2003016475-A2.

PD 27-FEB-2003.

PA (GEHO) GEN HOSPITAL CORP.
PA (KIRI) KIRIN BEER KK.
Percent Similarity: 43.02%
Best Local Similarity: 32.56%
 PD 07-MAR-2002.
PA (EPIG-) EPIGENOMICS AG.
Percent Similarity: 47.92%
Query Match: 14.39%
 43.06%
30.56%
14.39%
 43.06%
30.56%
14.39%
 ABQ53248 standard;
 Best Local Similarity:
 ID ADB06107 standar
DB Alloiococcus oti
PD 402003048304-A2.
PD 12-JUN-2003.
PA (AMHP) WYETH HO
Percent Similarity:
 US6551795-B1.
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Percent Similarity:
 ABQ32519 standard; DB Conservative: 11

ABQ32519 standard; DNA; 1246 BP.

DB Oligonucleotide for detecting cytosine methylation SEQ ID NO 19110.

PA (EPIG-) EBIGENORICS AG.

PA (EPIG-) EBIGENORICS AG.

PESTGORISHIARITY: 24.56$ Mismatcher.

RESULT 411

ID ABQ32518 standard; DNA; 1."

DE Oligonucleotide for PR SECONS PA (EPIG-) PR SECONS PA (EPIG-) PR
 ID ADO15235 standard; DNA, 435 BP.

DE Influenza A virus A/Moscow/10/99 (H3N2) partial nucleoprotein DNA.

PN WOO04041866-Al.

PD 21-MAY-2004.

PA (ADJU-) ADJUVANTIX LTD.

Percent Similarity: 42.86$ Conservative: 10

Best Local Similarity: 31.87$ Mismatches: 28

Ouery Match: Indels: 24
 ID ABX7212 standard; DNA; 75798 BP.

DE Mouse uridine diphosphphoglucuronosyl transferase gene locus.

PN WO20082897-A1.

PD 24-OCT-2002.

PA (GENE-) GENE STREAM PTY LTD.

PA (GENE-) GENE STREAM PTY LTD.

Conservative: 11

Best Local Similarity: 29.51% Mismatches: 7

Indels: 7
 ID NO:6651
 9
17
18
 10
33
6
 9
26
15
 15
32
19
 AAS77275 standard; cDNA; 874 BP.
DNA encoding novel human diagnostic protein #13079.
WO200175067-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 PD 23-AUG-2001.

PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Percent Similarity: 42.62% Conservative:

Best Local Similarity: 27.87% Mismatches:

Query Match:

RESULT 408
 Conservative:
Mismatches:
 DE AD12064 standard; DNA; 1754382 BP.

DE Alloicoccus otitis entire genome sequence SEQ PN W20203404-A2.

PA (AMHP) WYETH HOLDINGS CORP.

PACENT Similarity: 43.06$ COBERVATIVE: Best Local Similarity: 14.39$ Indels:
 ABV58775 standard; cDNA; 529 BP.
Human prostate expression marker cDNA 58766.
WO200160860-A2.
 Indels:
 PD 11-0CT-2001.

PA (HYSE-) HYSEQ INC.

Percent Similarity: 45.83%

Query Match: 14.29%
 45.16%
29.03%
14.39%
 PA (FARB) BAYER AG.
Percent Similarity:
Best Local Similarity:
 Query Match:
RESULT 406
 Query Match:
 RESULT 405
```

			ID NO 41371.			protein, Seq ID 1668.	ID NO: 7820. 0 0 15 15	ID NO: 2675.
16 27 0	16 44 44	9 29 10	SEQ 11 28 28 15	1 2 8 4 1 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	14 28 14		EQ I 10 25 28	SEQ
Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:	ene #2867. Conservative: Mismatches: Indels:	polynucleotide 6 Conservative: Mismatches: Indel8:	-2247. Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:	I novel secreted Conservative: Mismatches: Indels:	antigen DNA Si Conservative: Mismatches: Indels:	DNA fragment Conservative:
roent Similarity: 52.63% st Local Similarity: 24.56% sry Match: 14.29% SULT 412	ID ABA01231 standard; DNA; 1394 BP. DE T26F17.15 coding sequence. PN FR2806095-A1. PD 14-SEP-2001. PA (GENT-) GENTECH SARL. Percent Similarity: 36.89% Best Local Similarity: 23.77% Query Match: 41.29%	509 standard; DNA; 1880 BP. gillus fumigatus essential g 286090-A2 2002 ELITRA PHARM INC. milarity: 40.91% Similarity: 27.27% h:	DNA, 2685 BP. aster genomic 42.67% 28.00%	RESULT 415 ID AAQ40728 standard; DNA; 3045 BP. DE Human TPO gene lacking bases 2221- PN W09303146-A1. PD 18-FEB-1993. PA (RAPO/) RAPOPORT B. Percent Similarity: 46.15% Best Local Similarity: 28.21% Query Match: 14.29%	3 standard; DNA; 3072 PO gene. 46.Al. 1993. RAPOPORT B. 1arity: 46.15* imilarity: 14.29*	ID AAS26694 standard; DNA; 3659 BP. DE Human genomic DNA encoding partial PN W0200155322-A2. PD 02-AUG-2001. PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 36.90% Best Local Similarity: 25.00% Query Match:	RESULT 418 ID AL0512 standard; DNA; 3659 BP. DE Human reproductive system related PN W0200105320-A2. PD 02-AUG-2001. PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 36.90% Dest Local Similarity: 25.00% Query Match:	RESULT 419 ID ABL98023 standard; DNA; 3659 BP. DE Human testicular antigen encoding PN W0200155317-A2. PD 02-AUG-2001. PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 36.90\$

```
Human reproductive system related antigen DNA SEQ ID NO: 9679. WO200155320-A2.
 ABA18087 standard; DNA; 11599 BP.
Human nervous system related polynucleotide SEQ ID NO 1041B.
W0200159063-A2.
I-5-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 Human colon cancer antigen encoding cDNA SEQ ID NO:572 WO200122920-A2.
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23
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
Mismatches:
Indels:
 Mismatches:
 Query Match:
RESULT 425
ID ADH85055 standard; DNA; 744 BP.
DE Enteroccus faecalis polynucleotide #2940.
PN US6617156-B1.
 ABT19913 standard; DNA; 3880 BP.
Aspergillus fumigatus essential gene #2271.
WO200286090-A2.
 Indels:
 Indels:
 Indels:
 ADF82151 standard; DNA; 862 BP.
Leukaemia-related DNA sequence #2707.
 LOEKR. 1885.

LOEKR. DEUT KREBSFORSCHUNGSZENTRUM.

(UYLU-) UNIV LUDWIG MAXIMILIANS.

(HAFE/) HAFERLACH T.
 DNA; 32082 BP
 Human novel polynucleotide #871.
US2002132753-A1.
 AAH33516 standard; cDNA; 733 BP
 PN W0200122920-A2.
PD 05-APR-2001.
PA (HUMA) - HUMAN GENOME SCI INC.
Percent Similarity: 54 39%
Best Local Similarity: 29 82%
 PD 02-AUG-2001.
PD (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 38.61%
Best Local Similarity: 26.73%
 PN WOLVEL
PD 16-AUG-2001.
PA (HUMA-) HUMAN GENOME SC1
Percent Similarity: 53.33*
Best Local Similarity: 28.89*
North:
 DE Human novel polynuciecure.
PN US2002132753-A1.
PD 19-E8P-2002.
PA (ROSE/) ROSEN C A.
PA (RUBE/) RUBEN S M.
PA (BARA/) BARASH S C.
PA (BARA/) BARASH S C.
Bercent Similarity: 35.90%
Best Local Similarity: 25.00%
 44.23%
30.77%
14.18%
 54.39%
29.82%
14.18%
25.00%
14.29%
 PN W0200286090-A2.
PD 31-OCT-2002.
PD 31-CEITTA PHARM INC.
Percent Similarity: 40.91%
Best Local Similarity: 27.27%
 ID
DE EN.
DE EN.
PN US661.
PN US661.
PA (BOUC/) DOUL.
PA (BRSH/) BUSH D.
Parcent Similarity:
Local Similarity:
"Atch:
"At
 T 423
AAL06991 standard;
Sest Local Similarity:
 Best Local Similarity:
Query Match:
RESULT 427
 SCHOCH C.
KERN W.
 WO2003039443-A2.
 Percent Similarity:
 15-MAY-2003
 (SCHO/)
(KERN/)
 Query Match:
RESULT 420
 Query Match:
RESULT 424
 Query Match:
RESULT 422
 Query Matcl
RESULT 421
```

```
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:33755.
WO200157182-A2.
 AAS31096 standard; cDNA; 1106 BP.
Human diagnostic and therapeutic polynucleotide (DITHP) #111.
WO200162927-A2.
 10
33
15
 7 26
 14
23
3
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Mismatches:
Indels:
 Mismatches:
 Pseudomonas aeruginosa polynucleotide #16544.
US6551795-B1.
 Indels:
 Indels:
 Indels:
 NOVX related polynucleotide SEQ ID No 55.
WO200299062-A2.
 Novel human polypeptide NOV20b cDNA.
US2004018555-A1.
 PD 22-APR-2003.

PA (GENO-) GENOME THERAPEUTICS CORP.
Percent Similarity: 42.37*

Query Match: 14.18*
 ADO41689 standard; cDNA; 2482 BP
 ABD17940 standard; DNA; 1341 BP
 AAK78943 standard; DNA; 2052 BP
ID AAS31096 standard, c....,
DE Human diagnostic and therapeuti
PN W0200162927-A2.
PD 30-AUG-2001.
PA (INCY-) INCYTE GENOMICS INC.
Percent Similarity: 51.79%
Best Local Similarity: 35.17%
 PN MCCOLL.

09-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 54.39%

Best Local Similarity: 29.82%
 RESULT 431
ID ABT16044 standard; DNA; 2483
 PD 12-DEC-2002.
PA (CURA-) CURAGEN CORP.
Parcent Similarity: 40.74%
Best Local Similarity: 14.18%
 PA (ELLE, 40.73)
Percent Similarity: 40.739
Best Local Similarity: 28.408
14.188
 TAUDERR N. J. GUO X S. GUO X S. SHENOY S G. HJALT T. VOSS E Z. BOLDOG F L. MALYANKAR U M. PADIGARU M.
 RASTELLI L.
MACDOUGALL J R.
 LI L.
ZHONG M.
CASMAN S J.
GERLACH V.
SHIMKETS R A.
 PATTURAJAN M.
 GORMAN L.
PENA C E A.
KEKUDA R.
 SMITHSON G.
EDINGER S R.
 SPYTEK K A.
LEITE M W.
 ANDERSON D
ZERHUSEN B
 ELLERMAN K.
 MILLET
 JI W.
 29-JAN-2004.
(ANDE/) ANDE
 (HJAL/)
(VOSS/)
(BOLD/)
(MALY/)
(PADI/)
 Query Match:
RESULT 428
 Query Match:
RESULT 430
 (LILL/)
 (GERL/)
 (GORM/)
 (PATT/)
(SPYT/)
 (LEIT/)
(RAST/)
(MACD/)
(TAUP/)
(GUOX/)
(MILL/)
(SHEN/)
 ZERH/)
 (SMIT/)
 (EDIN/)
 Query Match:
RESULT 432
 (CASM/)
 KEKU/)
 PENA/)
 (MILL/)
 RESULT 429
```

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ID ACL.

DE Human oval.

PN WA201019281-A2.

PD 06-DEC-2001.

PA (CORI-) CORIA CORD.

PA (CORI-) CORIA CORD.

Percent Similarity: 57.58% Conselv.

Percent Similarity: 42.42% Mismatches: 0

RESULT 438

ID AAA02682 standard; cDNa; 509 BP.

DE Human colon cancer cell line polynucleotide sequence SEQ ID NO:2673.

PN WO9958675-A2.

19-NOV-1999.

THROW CORP.

Conservative: 5 Mismatches: 14 Mismatches: 10
 Hydrophobic domain protein cDNA HP10670 isolated from WERI-RB cells. WO200029448-A2.
 AAHS2705 standard; DNA; 1362 BP.
S. epidermidis open reading frame nucleotide sequence SEQ ID NO:803.
WO200134809-A2.
 ABN91304 standard; DNA; 1386 BP.
Staphylococcus epidermidis ORP nucleic acid sequence SEQ ID NO:767.
US6380370-B1.
 9
20
 10
19
 240
 (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY. (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO. It Similarity: 57.58 Conservative: Local Similarity: 42.428 Mismatches:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Mismatches:
Indels:
 Streptococcus polynucleotide SEQ ID NO 1523
WO200234771-A2.
 Indels:
 Prokaryotic essential gene #28391.

W20027183-A2.

03-OCT-2002.

(ELIT-) ELITRA PHARM INC.
 ID ABN91304 standard; DNA; 1386 BP.
DE Staphylococcus epidermidis ORF nuc
PN US6380370-B1.
PD 30-APR-2002.
PA (GENO-) GENOME THERAPEUTICS CORP.
Percent Similarity: 45.90%
 DNA; 1296 BP
 ADC87324 standard; DNA; 1300 BP Human GPCR gene SEQ ID NO:1777. EP1270724-A2.
 Query Match: 14.07%
RESULT 442
ID ACA46734 standard; DNA; 1359 BP
 ID A-
DE Strepto
PN WO2003471-A-
PN WO2003471-A-
PD 02-MAY-2002.
PA (CHIR-) CHIRON SPA.
Percent Similarity: 42.19$
Rest Local Similarity: 28.12$
Match:
 PA (SAGA) SAGAMI CHEM RES CENT.
PA (PROT) PROTISGENE INC.
Percent Similarity: 57.58
Best Local Similarity: 42.42$
 57.58%
42.42%
14.07%
 PN 17-MAY-2001.
PD 17-MAY-2001.
PA (GLAX) GLAXO GROUP LTD.
Percent Similarity: 45.90%
Best Local Similarity: 29.51%
Best Local Similarity: 14.07%
 45.90%
29.51%
14.07%
 42.42%
 Percent Similarity:
Best Local Similarity:
Query Match:
RESULT 443
 Local Similarity:
 Percent Similarity:
 25-MAY-2000
 02-JAN-2003
 Query Match:
RESULT 441
 Best
 ABK63760 standard; cDNA; 3726 BP.
Rat sequence differentially expressed in response to a hepatotoxin #1667.
WO200210453-A2.
 10
33
 10
33
15
 11
27
30
 11
27
30
 11
27
30
 1716.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
 Mismatches:
Indels:
 rat gene SEQ ID No
 Mismatches:
 Indels:
 ABT16043 standard; DNA; 2588 BP.
NOVX related polynucleotide SEQ ID No 53.
WO200299062-A2.
 ADO41687 standard; cDNA; 2588 BP.
Novel human polypeptide NOV20a cDNA.
US2004018555-Al.
 Toxicity-related gene, SEQ ID 3376. WO2003064624-A2.
 ВР
 ADB58350 standard; DNA; 3726 BP
 DNA; 3726 | related ra
 ID ABK63760 standaru; c....,
DE Rat sequence differentially e
NW0200210453-A2.
PD 07-FEB-2002.
PA (GENE-) GENE LOGIC INC.
Percent Similarity: 38.71%
Best Local Similarity: 26.88%
 DE NOVEL human polypeptide NOVE
DE NOVEL human polypeptide NOVE
DE STANDE/ ANDERSON D W.
DE CANDE/ ANDERSON D W.
DE CANDE/ ANDERSON D W.
DE CANDE/ ANDERSON D W.
DE CASM/ CASWAN S. J.
DE CASM/ CASWAN J. J.
DE CASM/ CASWAN S. J.
DE CASMAN J. J. W.
DE CASMAN J. W
 ID AET16043 standard; DNA; 2586
DE NOVX related polynucleotide
PN W0200299052-A2.
PD 12-DEC-2002.
PA (CURA-) CURAGEN CORP.
Percent Similarity: 40.74%
Query Match: 14.18%
RESULT 433
 38.71%
26.88%
14.18%
 PA (GENE-) GENE LOGIC INC.
Percent Similarity: 38.71%
Best Local Similarity: 26.88%
Query Match: 14.18%
RESULT 436
 PN WO200295000-A2.
PD 28-NOV-2002.
PA (GENE-) GENE LOGIC INC.
Percent Similarity: 38.71.
Best Local Similarity: 26.88
Query Match: 14.18
 ABT42014 standard;
Toxicity modelling
WO200295000-A2.
 07-AUG-2003
 Query Match
RESULT 434
```

			cdna.					
19	0 4	и н о 4.	encoding e: 10 8	0 14	10 26 8	<b>i</b>	10 26 8	162.
Mismatches: Indels:	cDNA.  CO LTD.  COnservative:  Mismatches: Indels:	Conservative: Mismatches: Indels:	protein- nservativ smatches:	Conservative: Mismatches: Indels:	one HPEBT96. Conservative: Mismatches: Indels:	secreted protein	Conservative: Mismatches: Indels:	de seguence #9062
0.35	H44494 standard; cDNA; 1419 BP. Iman UGT-isoenzyme 28 encoding cl 11297045-A. SHAN-2001. SHAN-) SHANGHAI BORONG GENE DEV ( 5. Similarity: 57.8% cal Similarity: 42.42% fatch: 14.07%	RESULT 446  ID ABT08325 standard; DNA; 1463 BP. DE Human NOV8a gene sequence. PN WO200246408-A2. PD 13-UNA-2002. PD 13-UNA-2002. Percent Similarity: 57.58% Best Local Similarity: 42.42% Query Match: 14.07%	standa lithp int 7031-A2. 2002. INCYTE larity: imilarit	MESULT 448  ID ABT08326 standard; DNA; 1742 BP. DE Human NOV8b gene sequence. PN WO200246408-A2. PD 13-UNN-2002. PA (CURA-) CURAGEN CORP. PA (CURA-) Similarity: 57.58% Best Local Similarity: 42.42% Query Match: 14.07%	7 standard; DNA; 1849 BP. ecreted protein gene 37 cl 90-A1. 1999. HUMAN GENOME SCI INC. larity: 44.26% imilarity: 27.87%	; cDNA; 1849 BP. encoding human	PA (FLOR) FLORENCE K A.  PA (HUJJ/) HU J.  PA (YUGG/) YU G.  PA (YUGG/) YU G.  PA (JANA/) JUAN R D.  Percent Similarity: 44.26%  Best Local Similarity: 27.87%  Query March: 14.07%	ACF70595 standard; DNA; 1860 BP. Photorhabdus luminescens nucleotide WO200294867-A2. 28-NOV-2002. (INSP ) INST PASTEUR.
Best Ouery	RESULT ID AN DE HI PN CI PD 3(0 PA (0 Percent Best Lo	RESULT ID AL DE HO PD 13 PA (( Percent Best Lo Quest Lo RESULT	RESUL ID DE DN PD PP Percel Best J	KESULT 4 ID ABT DE Hun PN WOZ PD 13- PA (CT Percent Best Loc	KESUL ID DE DE PN PD PA Percer Best	RESULT ID A DE DE PN I PA PA P	rce st ery SUL	ID DE PN PD

```
ID AAP93826 standard; cDNA; 1988 BP.

BE Human cDNA encoding a membrane or secretory protein clone PSEC0149.

BY EP1067182-20.

PA (HELL-) HELLX RES INST.

Percent Similarity: 57.58% Conservative: 5

Best Local Similarity: 42.42% Mismatches: 14

Query Match: 14.07% Indels: 0

RESULT 455.
 18
 2 7 0
 270
 114
 5
14
0
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
PA (CNRS) CNRS CENT NAT RECH SCI.

Percent Similarity: 53.85% Conservative.

Best Local Similarity: 41.03% Mismatches:

Query Match: 14.07% Indels:

RESULT 452

DE Human NF-kB activating gene SEQ ID NO 84.

PD 11-JUL-2002.

PA (ASAH) ASAHI KASEI KOGYO KK.

Percent Similarity: 42.42% Mismatches:

Query Match: 14.07% Indels:

RESULT 453

ID ACC42296 standard; cDNA; 1898 BP.

DE Human MAD Kinase cascade activator #15 cDNA.
 CDNA encoding novel human secreted protein #99. US2003027132-A1.
 Mismatches:
Indels:
 Mismatches:
Indels:
 AAV34252 standard; DNA; 2003 BP.
Human secreted protein gene 41 clone HNTME13.
WO9839446-A2.
 ABQ91954 standard; cDNA; 2018 BP.
Human NF-kB activating gene SEQ ID NO 86.
11-JUL-2002.
 DE Human secreted protein gene 41 clone
BE Human secreted protein gene 41 clone
BN (HumA-) Human GENOME SCI INC.
Best Local Similarity: 57.58 % Mil
RESULT 456

LO (Local Similarity: 42.42 % Mil
RESULT 456

LO (Local Similarity: 42.42 % Mil
RESULT 456

LO (Local Similarity: 42.42 % Mil
RESULT 456

RESULT 457

RESULT 4
 DE Human MAP Kinase cascade activator |
PN W02003008599-A1.
PD 30-ADN-2003.
PA (ASAH) ASAHI KASEI KOGYO KK.
Percent Similarity: 57.58% CC
Best Local Similarity: 142.42% M
Cuery Match: 14.07% ID
RESULT 454
ID AAF93826 standard; CDNA; 1988 BP.
```

```
PA (FENG/) FENG P.
PA (TOUN/) YOUNG P E.
PA (GREE/) GREENG J M.
PA (FERR/) FERRIE A M.
PA (HUJJ/) HU J.
PA (FLOR/) FLORENCE K P.
PA (FLOR/) FLORENCE K P.
PA (EBNE/) EBNER R.
PA (EBNE/) BREWER L A.
PA (BREW/) BREWER L A.
PA (SHIY/) SHI Y.
PERCORT SIMILARILY:
BEST LOCAL SIMILARILY:
 YOUNG P. E. GREENE J. M. FERRIE A M. DUAN R.
 BREWER L A.
 PD 26-JUL-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
 Percent Similarity:
 Query Match:
RESULT 466
 Query Match:
RESULT 465
 Query Match:
 Human secreted protein-encoding gene 8 cDNA clone HLHDL62, SEQ ID NO:18.
WO200132837-A1.
10-MAY-2001.
 AAA62075 standard; DNA; 2022 BP.
Hydrophobic domain protein cDNA HP10670 isolated from WERI-RB cells.
WO200029448-A2.
 Human transmembrane sugar transporter BioHSTR-encoding cDNA.
CN1293249-A.
02-MAY-2001.
 14
 277
 240
 14
 ID ABQ77696 stanuard, compared transporter block.

DB Human transmembrane sugar transporter block.

DR (N1293249-A.

DO 2-MAY-2001.

PA (SHEN-) SHENGYUAN GENE DEV CO LTD SHANGHAI.

Percent Similarity: 57.58% Conservative:

Best Local Similarity: 42.42% Mismatches:

14.07% Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 ACD08065 standard; cDNA; 2070 BP. cDNA encoding novel human secreted protein #41. US2003027132-A1.
 Human secreted protein encoding DNA clone vo22 WO200055375-A1.
 Mismatches:
 Mismatches:
 AAV34194 standard; DNA; 2070 BP.
Human secreted protein gene 41 clone HNTME13
WO9839446-A2.
 Indels:
 PA (SHAN-) SHANGHAI BIORIGIN GENE DEV CO LTD.
Percent Similarity: 57.58 Conserve
Best Local Similarity: 42.42$ Mismatch
Query Match: 14.07$ Indels:
 Indels:
 Indels:
 Indels:
 T 462
AAD08198 standard; cDNA; 2053 BP.
 CDNA; 2027 BP
 ID AAA62075 standard; DWA, CULLINA Hydrophobic domain protein cDNA PN W200029448-A2.
PD 25-MAY-2000.
PA (SAGA) SAGAMI CHEM RES CENT.
PA (FROT-) PROTEGENE INC.
PErcent Similarity: 42.42%
Best Local Similarity: 42.42%
 AAC59822 standard; DNA; 2051 BP
 PN W09839446-A2.
PD 11-SEP-1998.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 57.58%
Best Local Similarity: 42.42%
 (HUMA-) HUMAN GENOME SCI INC.
 KOGYO KK.
 57.58%
42.42%
14.07%
 57.58%
42.42%
14.07%
 PD (ALPH-) ALPHAGENE INC.
Percent Similarity: 57.58%
Best Local Similarity: 42.42%
Ouery Match: 14.07%
 (ROBE/) ROBEN S M.
(ROSE/) ROSEN C A.
(FISC) FISCHER C L.
(SOPP/) SOPPET R C.
(CART/) CARTER K C.
(BEDN/) BEDNARIK D R.
(TUGG/) YU G.
(YUGG/) YI G.
PA (ASAH) ASAHI KASEI KC
Percent Similarity; 57.
Best Local Similarity: 42.
Query Match: 14
RESULT 458
 AAH20413 standard; cD
Human BioATPase cDNA.
WO200130836-A1.
 PA (HUMA-) HUMAN GENO)
Percent Similarity:
Best Local Similarity:
Query Match:
 06-FEB-2003.
 21-SEP-2000
 Ouery Match:
RESULT 464
ID ACD08065
DE CDM enco
PN US200302'
PD 06-FB-2
PA (ROSE/)
PA (SOPP/)
 Query Match:
RESULT 459
 RESULT 461
```

```
Human secreted protein-encoding gene 8 cDNA clone HDFQB93, SEQ ID NO:31.
WO200132837-A1.
 ABZ35234 standard, cDNA, 3170 BP.
Human gene expression profile polynucleotide SEQ ID NO 345.
WO200274979-A2.
 DE Human soft tissue sarcoma upregulated DNA - SEQ ID 7485.
PN W0200404838-A2.
PD 10-JUN-2004.
PA (PROT-) PROTBIN DESIGN LABS INC.
Percent Similarity: 57.58$ Mismatches: 5
Best Local Similarity: 14.42$ Mismatches: 14
Chery Match: 14.07$ Indels: 0
 ID 6085
 270
 140
 240
 10
42
11
 C. albicans specific gene, orf6.7416, DNA sequence. W02004056965-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 ADQ23265 standard; DNA; 2555 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ
WO2004048938-A2.
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Mismatches:
 Mismatches:
 Mismatches:
 Indels:
 Indels:
 Indels:
 DE Human polynucleotide SEQ ID NO 3720.
PN WO200153312-A1.
PD 26-JUL-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity: 57.58$ Conser Best Local Similarity: 42.42$ Mismack Query Match:
 Human polynucleotide SEQ ID NO 148.
WO20013312-Al.
 10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
 AAD08211 standard; cDNA; 2072 BP.
 ID AAD08211 standard; curr, coll by Human secreted protein-encoding by WOC0132837-A1.

PD 10-MAY-2001.

PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 57.58% Best Local Similarity: 14.07%
 RESULT 468
ID ADP98650 standard; DNA; 2355 BP
 DNA; 2551 BP
 PA (ELIT-) ELITRA PHARM INC.
PA (ELIT-) ELITRA CANADA LITD.
Percent Similarity: 36.21%
Best Local Similarity: 36.21%
Query Match: 14.07%
RESULT 469
ID ADQ24665 standard; DNA; 2551
 57.58%
42.42%
14.07%
 43.01%
32.26%
14.07%
 57.58%
42.42%
14.07%
HU J.
FLORENCE K A.
OLSEN H S.
EBNER R.
```

nce molecule (PMMM)-47 gene.

equence SEQ ID NO:4079.

PA (MILL-) MILLENIUM PHARM INC. Percent Similarity: 43.01% Mismatches: 10 Best Local Similarity: 32.26% Mismatches: 42 Ouery March: 14.07% Indels: 11 RESULT 480 ID ADF90653 standard, DNA, 3935 BP. DB Human hepatic-fibrosis disease marker SEQ ID 115. PD 16-SEP-2003.	PA (SUMU) SUMITOMO SELYAKU KK.  Percent Similarity: 43.01\$ Conservative: 10  Best Local Similarity: 43.05\$ Mismatches: 42  Query Match: 14.07\$ Indels: 11  RESULT 481  ID ADETYOG7 standard; DNA; 4025 BP.  DE Human protein modification and maintenance molecule (PMWM) -47  PN WO2003063688-A2.	PD 07-AUG-2003. PD (10.4 ) INCYTE GENOMICS INC. PA (INCY-) INCYTE GENOMICS INC. Percent Similarity: 43.01% Mismatches: 42 Query Match: 14.07% Indels: 11 RESULT 482 ID AAHS415 standard; DNA, 4095 BP. DE S. epidermidis genomic polynucleotide sequence SEQ ID NO:4075 PD 17-MAY-2001	(GLAX ) GLAXO GROUP LTD. cent Similarity: 45,90% st Local Similarity: 29,51% sty Match: 14.07% sulr 483	ABA16110 standard; DNA; 4678 B Human nervous system related pw0200159063.A2. 16-AUG-2001. (HUMA-) HUMAN GENOME SCI INC. coent Similarity; 47.06% st Local Similarity; 32.35% sry Match:	ABOS4837 standard; cDNA; 5169 Human ovarian antigen HNOKM38 W0200200677-A1. 03-JAN-2002. (HUMA-) HUMAN GENOME SCI INC. cent Similarity: 44.26 \$ st Local Similarity: 27.87 \$ sty Match: 14.07 \$ sty Match: 14.07 \$ sty Match: 2523 \$ ABF74193 standard; cDNA; 5223	/hippocampus DNA KENKYUSH KPRESS KK. 46.03# 31.75# 14.07# ; cDNA; 5387	gry st.
PD 26-SEP-2002.  PA (ORTH ) ORTHO CLINICAL DIAGNOSTICS INC.  Percent Similarity: 43.01% Conservative: 10  Best Local Similarity: 32.26% Mismatches: 42  Query Match: 14.07% Indels: 11  RESULT 472  ID ADE79065 standard; DNA; 3273 BP.  DB Human protein modification and maintenance molecule (PMMM) -45 gene.  PN WC20030636588-20	rce st sry sur	MOZDO20677-AL WO2D020677-AL 03-JAN-2002. (HUMA-) HUMAN GENOME SCI INC. CCERT SIMILATITY: 32.26\$ Mismatches: st Local Similarity: 32.26\$ Mismatches: TY MATCH: 14.07\$ Indels: SULT 474 AAH54886 standard; DNA; 3553 BP.	rce st	RESULT 475  ID AAT97610 standard; cDNA; 3854 BP.  DE Human E2A-binding protein cDNA.  PN W0973900-A1.  PD 18-SEP-1997.  PA (HARD) HARVARD COLLEGE.  Percent Similarity: 43.01% Mismatches: 10  Best Local Similarity: 14.07% Indels: 11	RESULT 476  ID ADE79063 standard; DNA; 3872 BP.  DE Human protein modification and maintenance molecule (PMMM) -43 gene.  PN WC2003063688-A2.  PN G2003063688-A2.  PA G1NCY-) INCYTE GENOMICS INC.  PACTOR Similarity: 43.01% Mismatches: 42  Query Match: 14.07% Indels: 11  RESULT 477	DE Human cDNA differentially expressed in lung cancer #133.  DE Human cDNA differentially expressed in lung cancer #133.  PN US2003065157-A1.  PD 03-APR-2003.  PA (LASE/) LASEK A W.  Percent Similarity: 32.26* Mismatches: 42  Query Match: 14.07* Indels: 11  RESULT 478  ID ABZ34770 standard; cDNA; 3935 BP.  DE Coding sequence SEQ ID 128, upregulated in osteogenesis.  PN WOODD S1745-AC.	SUL SUL

Conservative: Mismatches: Indels: GCONSERVATIVE: Mismatches: Indels: Indels: Mismatches: Indels: Mismatches: Indels:	INCYTE CORP.  larity: 44.26* Conservative: 10  imilarity: 27.87* Mismatches: 26  48.84 Standard, DNA; 5886 BP.  coff tissue sarcoma-upregulated DNA - SEQ ID 835  48.85 BP.  coff tissue sarcoma-upregulated DNA - SEQ ID 835  48.26 Mismatches: 26  imilarity: 27.87* Mismatches: 86  estandard, cDNA; 5887 BP.  ERISTOL-WYERS SQUIBS CO.  14.07* Indels: 86  ERISTOL-WYERS SQUIBS CO.  2003.  ERISTOL-WYERS SQUIBS CO.  AMILIARIUM PREDICTIVE MEDICINE INC.  14.07* Indels: 8  14.07* Mismatches: 26  imilarity: 27.87* Mismatches: 26  imilarity: 27.87* Mismatches: 26  imilarity: 27.87* Mismatches: 26  Indels: 8  7 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  6 standard, cDNA; 5893 BP.  7 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  7 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  7 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  7 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  6 standard, cDNA; 5993 BP.  7 standard, cDNA; 5993 BP.  8 mismatches: 8  8 mismatches: 9  8 mismatches	PD 02-A PA (HUM Percent S Best Loca	QUELY MAC RESULT 49 ID ABX6	ላያ ተያ	Percent S Best Local Query Math	1.0 DB DB PN PD PD	Percent S Best Loca Quert Coca RESULT 49 ID AAD3	DE PN PD PD	Percent S. Best Loca. Query Math	DD DE PN PD	Percent S. Best Loca: Query Mate	ID AAC80 DB Comp. PN W0.20 PD 28-01 PD 28-01 PA (PHAC	PA (WID PACENT S Best Loca Query Mat	Percent St. Percent St. Pert Local Query Matc	Percent Si Best Local Query Marc RESULT 503	DE HUMAN PN EP127 PD 02-07 PA (NAAI PA (ADSC
The property of the property o	0RP. 44.26%   127.87%   14.07%   6 barcoma-upregg   14.07%   14.07%   14.07%   27.87%   14.07%   27.87%   14.07%   27.87%   14.07%   27.87%   14.07%   37.893 BE   27.87%   14.07%   40, CDNA, 5893 BE   27.87%   14.07%   41.07%   42.26%   14.07%   27.87%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%   14.07%	10 26 8	83	10 26 8		10 26 8		10 26 8		10 26 8	543	10 26 8		8 17 4	ence.	9 25 25
(1) 10 11 11 11 11 11 11 11 11 11 11 11 11	ORP.  14.26%  27.87%  24.26%  e sarcoma-upregr  DESIGN LABS INC  27.87%  14.07%  d, cDNA, 5887 Bi  ker polynucleot:  27.87%  14.07%  d, cDNA, 5893 Bi  At.26%  27.87%  14.07%  d, cDNA, 5903 Bp.  At.26%  27.87%  14.07%  d, cDNA, 5903 Bp.  ESTCOMA-upregu  UM PREDICTIVE ME  27.87%  14.07%  d, cDNA, 5903 Bp.  ESTCOMA-upregu  DESIGN LABS INC.  27.87%  14.07%  d, DNA, 9345 Bp.  id synthase gene  TECHNOLOGY LLC.  14.07%  d, DNA, 12017 Bi  d, DNA, 12017 Bi  sporter protein  K.  K.  K.  K.  K.	Conservative: Mismatches: Indels:	DNA -	Conservative: Mismatches: Indels:	SEQ ID NO:4	Conservative: Mismatches: Indels:	CDNA 26860.	tive es:	CDNA 21018.	tive es:	SEQ	a		Conservative: Mismatches: Indels:	enomic DNA sequ	Ó
	y-) INCYTE CORBINITY:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 PROTEIN DES  18 Similarity:  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  11 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 PROTEIN DES  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  10 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 PROTEIN DES  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 Similarity:  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  10 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 Similarity:  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  10 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 Similarity:  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  10 Similarity:  11 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 Similarity:  18 Similarity:  18 Similarity:  19 Similarity:  10 Similarity:  10 Similarity:  11 Similarity:  11 Similarity:  12 Similarity:  13 Similarity:  14 Similarity:  15 Similarity:  16 Similarity:  17 Similarity:  18 Simi	24.26% 27.87% 14.07%	DNA; 5886 BP.	51GN LABS INC. 44.26% 27.87% 14.07%	cDNA; 5887 BP. : polynucleotic	TRS SQUIBB CO. 44.26% 27.87% 14.07%	cDNA; 5893 BP.	PREDICTIVE MEL 44.26% 27.87% 14.07%	cDNA; 5893 BP. ression marker	PREDICTIVE MED 44.26% 27.87% 14.07%	DNA; 5903 BP. sarcoma-upregul		9345 hase		ដូដ	l. 37.93\$ 27.59\$ 14.07\$

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- SEQ ID 3698.
 60321 standard, cDNA, 16100 BP.
A encoding novel human musculoskeletal system antigen #2665.
002147140-A1.
 36071 standard, DNA, 16161 BP.
an adipocyte enhancer binding protein encoding genomic DNA.
00224741-A2.
 31071 standard; DNA; 16100 BP.
an musculoskeletal system-associated genomic DNA
004009488-A1.
 8
70
70
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38
 142
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15
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34
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34
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 JAN 2003.
AD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
SC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
SImilarity: 44.44% Conservative: 8
al Similarity: 33.33% Mismatches: 2
tch: I4.07% Indels: 2
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Indels:
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Mismatches:
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Mismatches:
Indels:
 03
87620 standard; DNA; 144792 BP.
270724-A2.
 teriophage 77 complete genome sequence.
00032825-A2.
 ACG6106 standard; CDNA; 41708 BP.
COMPLETE genome of bacteriophage 77.
CO20146583-A2.
PHAG-30-1.
PHAG-) PHAGETECH INC.
WILL, WILLIAMS K M.
CSIMILARILY: 43.55$ CONSTOCAL Similarity: 27.42$ Missm COAL Similarity: 14.07$ Inde:
 58247 standard; DNA; 41708 BP.
 JOHN-2004.

JUAN-2004.

JUAN-2004.

Similarity: 38.52%

Occal Similarity: 24.59%
MA-) HUMAN GENOME SCI INC.
Similarity: 38.52%
al Similarity: 24.59%
tch:
 COSE) ROSEN C A.
ROSEN S M.
SARA/) BARASH S C.
Similarity: 38.52%
latch: milarity: 24.59%
 43.01%
32.26%
14.07%
 AG-) PHAGETECH INC.
Similarity: 43.55%
al Similarity: 27.42%
tch:
 50.67%
36.00%
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 -MAR-2002.
YAN/) RYAN J W.
Similarity:
cal Similarity:
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cal Similarity:
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 TUN-2000
 RESULT
ID AL
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ADC87619 standard; DNA; 349981 BP

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D ABZ13684 standard; DNA; 1347 BP.
DE Arabidopsis thaliana stress regulated gene SEQ ID NO 1489.
DE Arabidopsis thaliana stress regulated gene SEQ ID NO 1489.
PN WOZOO16655-A2.
PA (SCRI) SCRIPPS RES INST.
PA (SCRI) STNGENTA PARTICIPATIONS AG.
PERCENT Similarity: 45.83% Conservative: 8
Best Local Similarity: 34.72% Mismatches: 26
Query Match: 13.96% Indels: 13
RESULT S16
ID AAC39127 standard; DNA; 1535 BP.
DE Arabidopsis thaliana DNA; Iragment SEQ ID NO: 23472.
 AAF80403 standard; cDNA; 1707 BP.
Nuclectide sequence of a plant transcription factor G748.
WO200126459-A2.
 Arabidopsis thaliana transcription factor G748 cDNA W0200136444-A1.
 ACF72008 standard; DNA; 1272 BP.
Photorhabdus luminescens nucleotide sequence #10475.
WO200294867-A2.
 8
26
13
 10 26
 AAC51442 standard; DNA; 1347 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 68538.
EP1033405-A2.
 DNA encoding Acinetobacter baumannii protein #434.
US6562958-B1.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Conservative:
Mismatches:
 Conservative:
Mismatches:
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 25-MAY-2001.
(MEND-) MENDEL BIOTECHNOLOGY INC.
(RIEC/) RIECHMANN J L.
(REUB/) REUBER L.
 PA (GENO-) GENOME THERAPEUTICS CORP. Percent Similarity: 50.91%
 (MEND-) MENDEL BIOTECHNOLOGY INC
 CDNA; 1707 BP
 BP
 (INSP) INST PASTEUR.
(CNRS) CNRS CENT NAT RECH SCI.
 ADA29147 standard; DNA; 1068
 45.83%
34.72%
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13.96%
 Percent Similarity: 50.91%
Best Local Similarity: 32.73%
Query Match: 13.96%
 Percent Similarity: 47.00%
Best Local Similarity: 26.00%
Query Match: 13.96%
50.91%
32.73%
13.96%
 RIECHMANN J L
 ID AAC39127 standard; D
DE Arabidopsis thaliana
PN EP1033405-A2.
PD 06-SEP-2000.
Percent Similarity: 4
Best Local Similarity: 3
 SAMAHA R.
CREELMAN R.
KEDDIE J.
 RATCLIFFE O.
 AAD05834 standard;
 Best Local Similarity:
Query Match:
RESULT 515
 Best Local Similarity:
 Best Local Similarity:
Query Match:
 REUBER L
 HEARD J.
 JIANG C.
 ID AAF80403 standa
DE Nucleotide seque
PD 19-APR-2001.
PA (RATC) RATCLIF
PA (RATC) RATCLIF
PA (RATC) RATCLIF
PA (RAEAR) SAMAHA
PA (CREB) CREBLAB
PA (CREB) CREBLAB
PA (CRED) REDDIE
PA (TRED) REDDIE
PA (TRED) REDDIE
PA (REUM) JIANG C
PA (REUM) RECHABER
PA (REUM) RECHABER
PA (RECHABER
PA (RECH
 Percent Similarity:
 Percent Similarity:
 06-SEP-2000
 13-MAY-2003
 Query Match:
 Query Match:
RESULT 518
 Query Match:
RESULT 514
 RESULT 517
 RESULT 513
 ID ADK56099 standard; DNA; 644 BP.

ID Plant DNA sequence which confers altered metabolic characteristic #3482.

BN W02003020936-A1.

PD 13-MAR-2003.

PA (DOWC) DOW CHEM CO.

PA (DOWC) DOW AGROSCIENCES LLC.

Percent Similarity: 45.83 Mismatches: 26

Query Match: 13.96% Indels: 13
 ADK59802 standard; DNA; 679 BP.
Plant DNA sequence which confers altered metabolic characteristic #7185.
WO2003020936-A1.
13-MAR-2003.
 Bovine EST associated with lactation/muscle/fat deposition #11206.
US2002137139-A1.
26-SEP-2002.
 33
 33
 272
 ABZ18469 standard; cDNA; 386 BP.
Group III cDNA cancer related clone SEQ ID NO:895.
W0200278516-A2.
(CORI-) CORIXA CORP.
 DE Human GPCR related policy.

PN EP1270724-A2.

PD 02-JAN-2002.

PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.

PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

Percent Similarity: 44.44% Conservative:

Best Local Similarity: 13.33% Mismatches:

14.07% Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
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 Ouery Match:
RESULT 510
ID ADG32540 standard; DNA; 879 BP.
ID BAR encoding the human Claudin-23 protein.
PN WO2003100001-A2.
PD 04-DEC-2003.
PA (IMWV) IMMUNEX CORP.
Percent Similarity: 41.18% Mismatches
Described Similarity: 130.88% Mismatches
 Mismatches:
Indels:
 Mismatches:
Indels:
 RESULT 507
ID ACD05669 standard; cDNA; 630 BP.
DE cDNA encoding novel human polypeptide #199.
PN WO200323013-A2.
 Prokaryotic essential gene #2767.
03-0007183-A2.
 T 505
ABX46041 standard; cDNA; 383 BP
 (DOWC) DOW CHEM CO. (DOWC) DOW AGROSCIENCES LLC.
 PD 20-MAR-2003.

PD 20-MAR-2003.

PErcent Similarity: 41.18%
Best Local Similarity: 30.88%
 Percent Similarity: 45.83%
Best Local Similarity: 34.72%
Query Match: 13.96%
 Percent Similarity: 50.00%
Best Local Similarity: 38.46%
Query Match: 13.96%
 44.62%
 Best Local Similarity: 32.31%
 (ELIT-) ELITRA PHARM INC
 (BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
 ID ABX46041 standard
DB Bovine EST assoc
DB CO02131139-A1.
PD 26-SEP-2002.
PA (BYAT/) BYATT J (PAT/) PATT (PAT/) PATT (PAT/) PA (MATH/) MATHALABA (MARK/) WARREN W
PETCENT Similarity:
 Query Match:
RESULT 509
 Query Match:
RESULT 511
 Query Match:
RESULT 506
 Query Match:
 Query Match:
RESULT 508
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us-09-989-293a-377.rng.spdi

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ABK65211 standard; cDNA; 1707 BP.
Arabidopsis cDNA encoding a transcription factor #63
28-FBB-2002.
 AAD47515 standard; DNA; 1707 BP.
Arabidopsis thaliana G748 transcription factor DNA.
WO200274917-A2.
PA (KEDD/) KEDDIE J.
PA (HRAK) RATCLIFFE O.
PA (SAMALA) SAMAHA R.
PETCENT Similarity: 34.72% Mismatches:
Query Match: 13.96% Indels:
RESULT S19
DE ARabidopsis cDNA encoding a transcription factor MOZOJS675-A1.
PA (MEND-) BENDEL BIOTECHNOLOGY INC.
PA (NEND-) PRENDEL BIOTECHNOLOGY INC.
PA (REDD/) KEEDIE J.
PA (HEAR) HEARD J.
PA (HEAR) HEARD J.
PA (REDD/) KEEDIE J.
PA (REDD/) KEEDIE J.
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Mismatches:
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Mismatches:
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 Mismatches:
Indels:
 Mismatches:
 Plant transcription factor polynucleotide #78 US2004019927-A1.
 Indels:
 ID AAD47515 standard; DNA; 1707 BP.
DE Arabidopsis thaliana G748 transcript
PN W0200274917-A2.
PD 26-SEP-2002.
PA (MEND-) MENDEL BIOTECHNOLOGY INC.
Percent Similarity: 45.83%
Cuery Match:
RESULT 523
 20-FEB-2003. (MEND-) MENDEL BIOTECHNOLOGY INC.
 ADI41692 standard; DNA; 1707 BP.
 1D ADE31478 standard; cDNA; 170
DE Plant yield related polynuc.
PN WC200301228-A2.
PD 20-FEB-2003.
PA (WEND-) MENDEL BIOTECHNOLOGY
Percent Similarity: 45.83%
Best Local Similarity: 34.72%
Query Match:
RESULT 522
 29-JAN-2004.
SHER/) SHERMAN B K.
(RIEC/) RIECHMANN J L.
(JIAN/) JIANG C.
(HEAR/) HEARD J B.
(HAAK/) HAAKE V.
(CREC/) CREELMAN R A.
(RATC/) RATCLIFFE O.
(ADAM/) ADAM L J.
(REUB/) REUBER T L.
(KEDD/) KEDDIE J.
(REUG/) BROUN P E.
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ADE07093 standard; DNA; 1716 BP.
Novel coding sequence (useful for identifying genetic disorders) #159.
WO2003054152-A2.
 8
26
13
 8
26
13
 14
28
13
 33
 33
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
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Mismatches:
 Mismatches:
Indels:
PA (PILG/) PILGRIM M L.
PA (DUBE/) DUBELL A N.
PA (PINGS) YU G.
PA (PINGS) YU G.
PA (WINGS) YU G.
Percent Similarity: 45.83% Conservativ
Best Local Similarity: 13.96% Mismatches:
Query Match: 13.96% Indels:
RESULT 5.24
ID AD002084 standard; CDNA; 1707 BP.
DE Thalecress transcription factor cDNA #249.
PA (ZHAN/) ZHANG J.
PA (ZHAN/) ZHANG J.
PA (ZHAN/) ZHANG J.
PA (HEAR/) HEARD J E.
PA (HEAR/) HEARD J E.
PA (HEAR/) HEARD J E.
PA (RIEC/) RIECHMANN J L.
PA (RIEC/) RECHMANN J L.
PA (RIEC/) REUBER T L.
PA (KEDD/) KEDDIE J S.
PA (KEDD/) KEDDIE J S.
PA (CRED/) YU G.
PA (CRED/) YU G.
PA (CRED/) NEUBER T L.
PA (CRED/) REDIE J S.
PA (CRED/) REMBER N L.
PA (CRED/) REMBER N R.
PA (CRED/) REMBER N R.
PA (CRED/) REMBERNAN B K.
PA (CREE/) CREELMAN R A.
PA (CREE/) CREELMAN B K.
PECCENT SIMILATICY: 13.90% TARACTIFES O.
PA (KNATC) RATCLIFFE O.
PA (KNATC) R
 Indels:
 ABTI6021 standard; DNA; 1719 BP.
NOVX related polynucleotide SEQ ID No 9.
WO200299062-A2.
 ADO41643 standard; cDNA; 1719 BP.
Novel human polypeptide NOV5a cDNA
US2004018555-A1.
 Prokaryotic essential gene #2142.
WO200277183-A2.
 D ACA20485 standaru, L.M., -
DE Prokaryotic essential gene #2
DN W020027183-A2.
DD 03-OCT-2002.
PA (ELIT-) ELITRA PHARM INC.
Percent Similarity: 44.59$
Best Local Similarity: 25.68$
 ACA20485 standard; DNA; 1713
 D ABT16021 standard; DAR, CONTROL OF NOVX related polynucleotide S PN WO200299062-A2.

PD 12-DEC-2002.

PA (CURA-) CURAGEN CORP.

Percent Similarity: 41.18%

Best Local Similarity: 30.88%
 PA (HYSE-) HYSEQ INC.
Percent Similarity: 41.18*
Best Local Similarity: 30.88*
Query Match: 13.96*
RESULT 527
 nt Similarity: 45.83%
Local Similarity: 34.72%
Match: 13.96%
 29-JAN-2004.

(ANDE/) ANDERSON D W.
(ZERH/) ZERHUSEN B D.
(LILLL.) LI L.
(ZHON/) ZHONG M.
(CASM/) CASMAN S J.
 LI L.
ZHONG M.
CASMAN S J.
GERLACH V.
SHIMKETS R A.
GORMAN L.
 03-JUL-2003.
(HYSE-) HYSEQ INC.
 Query Match:
RESULT 526
 (GERL/)
(SHIM/)
(GORM/)
 Query M
RESULT
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```
PD 02-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 45.31%
 Query Match:
RESULT 535
 Query Match:
 Query Match:
 Query Match:
RESULT 540
 Human cardiovascular system antigen genomic DNA SEQ ID No 2110.
WO200155321-A2.
 Human reproductive system related antigen DNA SEQ ID NO: 8995.
WO200155320-A2.
 8479
 õ
 AAS30529 standard; DNA; 10812 BP.
DNA encoding novel prostate gland antigen, Seq ID No 387.
WO200155447-A1.
 SEQ ID
 12
27
3
 ABA00966 standard; cDNA; 2052 BP.
CGDD-7 coding sequence, Incyte ID No. 1820882CB1.
WO2003008553-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 ABL19002 standard; DNA; 21231 BP.
Drosophila melanogaster genomic polynucleotide
 Mismatches:
Indels:
 Mismatches:
Indels:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Indels:
 Indels:
 AAS36610 standard; DNA; 22008 BP.
 AAL06307 standard; DNA; 10812 BP
 DE Humau.
PN WO200155320-Az.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 46.94%
Best Local Similarity: 32.65%
13.96%
 BP
 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 ADH69503 standard; DNA; 6251
 Rat dopamine D1 receptor DNA US2003170741-A1.
 PN WC2003008553-A2.
PD 30-ADN-2003.
PA (INCY-) INCYTE GENOMICS INC.
Percent Similarity: 41.18%
 (PROC) PROCIER & GAMBLE CO.
 PA (PEAGE / - Percent Similarity: 49,12%
Best Local Similarity: 28,81%
13.96%
 Percent Similarity: 46.94%
Best Local Similarity: 32.65%
Query Match: 13.96%
 Best Local Similarity: 41.18% Query Match: 13.96% RESULT 529
 Percent Similarity: 41.18%
Best Local Similarity: 30.88%
Query Match: 13.96%
 nilarity: 50.94%
Similarity: 37.74%
h: 13.96%
 HJALT T.
VOSS E Z.
BOLDOG F L.
MALYANKAR U M.
PADIGARU M.
 RASTELLI L.
MACDOUGALL J R.
 PENA C E A.
KEKUDA R.
PATTURAJAN M.
 EDINGER S R.
MILLET I.
ELLERMAN K.
 AUPIER R J.
 GUO X S.
MILLER C E.
SHENOY S G.
 SPYTEK K A.
 (PEKE) PE CORP NY
 SMITHSON G.
 T 533
ABL19002 standard;
 LEITE M W.
 WO200171042-A2.
 Percent Similarity:
 JI W.
 11-SEP-2003
 (SMIT/)
(EDIN/)
(MILL/)
(ELLE/)
 Query Match:
RESULT 530
 Query Match:
RESULT 532
 Query Match:
 (TAUP/)
(GUOX/)
(MILL/)
(SHEN/)
 (HJAL/)
(VOSS/)
(BOLD/)
(MALY/)
 (KEKU/)
 (SPYT/)
(LEIT/)
(RAST/)
(MACD/)
 PADI/)
 JIWW/)
 RESULT 531
 PA (1)
PA (2)
PA (1)
 Best
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AAK83212 standard; DNA; 68356 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:38024.
WO200157182-A2.
09-AUG-2001.
 Secreted protein gene 33 genomic fragment HBJAB02, SEQ ID NO:930.WO200277013-A2.
 No 242
 8170
 Human secreted protein encoding genomic DNA SEQ ID NO 900.
WO200277186-A2.
 Human nervous system related polynucleotide SEQ 1D NO WO200159063-A2.
 - SEQ ID
 RESULT 539
ID ADEFT704 standard; DNA; 22008 BP.
DB Human cardiovascular system related genomic DNA #870.
PN US2003059908-A1.
 9
18
17
 9
17
 118
 Conservative:
Mismatches:
Indels:
 ADA98459 standard; DNA; 22008 BP.
Human secreted protein-related DNA sequence #52.
WO2003004623-A2.
 Conservative:
 Human secreted protein-related DNA sequence #45.
WO200292787-A2.
21-NOV-2002.
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
Mismatches:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Mismatches:
 Mismatches:
Indels:
 Human secreted protein-related DNA sequence WO200277188-A2.
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 RESULT 541
ID ABZ67377 standard; DNA; 22008 BP.
 T 538
ADC20627 standard, DNA; 22008 BP
 ABT16888 standard; DNA; 22008 BP
 RESULT 536
ID ABZ73783 standard; DNA; 22008 BP
 ABA15839 standard; DNA; 22008 BP
 03-OCT-2002. (HUMA-) HUMAN GENOME SCI INC.
 PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 45.31%
 PN MCACCACOLO.
PD 16-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 45.31%
Best Local Similarity: 31.25%
 PD 16-JAN-2003.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 45.31%
Best Local Similarity: 31.25%
 (HUMA-) HUMAN GENOME SCI INC
 45.31%
31.25%
13.96%
 Percent Similarity: 45.31%
Best Local Similarity: 31.25%
Query Match: 13.96%
 Percent Similarity: 45.31%
Best Local Similarity: 31.25%
Query Match: 13.96%
Percent Similarity: 45.31%
Best Local Similarity: 31.25%
Ouery Match: 13.96%
 13.96%
 Best Local Similarity:
 03-OCT-2002
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PD 23-AUG-2001. PA (MILL-) MILLENNIUM PRED Percent Similarity: 45.0 Best Local Similarity: 31.8 Onery Match:	0 standard; rostate expr	PN WCZULEUGBOL-AZ. PD AUG-2001. PA (MILL-) MILLENNIUM PRED Percent Similarity: 45.0	nry Match: SULT 553 ABK45654 standard; CDNA encoding color	COR	Deer Local Similarity: 26.8 Query Match: 13.8 RESULT 554 ID ABV33703 standard; CDNA DE Human prostate expressi	PN WO200160860-A2. PD 23-AUG-2001. PA (KILL-) MILLENNIUM PRED: Percent Similarity: 45.01	Best Local Similarity: 31.8' Query Match: 13.8' RESULT 555 ID ACC76835 standard: DNA;	MOZO 03020905-A2. 13-MAR-2003. (DOWC ) DOW CHEM (		·	Dest Bocal Similarity: 25.3 Query Match: 13.89 RESULT 557 ID MRV42605 standard: CDNA	H	. ÷ =	PD 28-SEP-2000 PD 28-SEP-2000 PA (NOVO) NOVO NORDISK BIG Percent Similarity: 43.72 Best Local Similarity: 28.75 Query Match:	rasol, 359 ID Ango9341 standard; DNA; DE Listeria innocua DNA sec PN W0200228891-A2.
Conservative: 12 Mismatches: 34 Indels: 77	gen genomic sequence SEQ ID NO:22095.	Conservative: 12 Mismatches: 34 Indels: 77	Conservative: 21 Mismatches: 26 Indels: 27	Conservative: 21 Mismatches: 26 Indels: 27	Ġ.	Conservative: 13 Mismatches: 32 Indels: 23	p. rphic nucleotide #382.	Conservative: 23 Mismatches: 24 Indels: 18	P. 1-30F3 SEQ ID NO:21.	Conservative: 17 Mismatches: 27 Indels: 23	BP, sed in granulocytic cells #920.	Conservative: 5 Mismatches: 18 Indels: 7	tion/muscle/fat deposition #11197.	Conservative: 13 Mismatches: 26 Indels: 12	cDNA 12560.
50 m E	ID AAK67283 standard; DNA; 68356 BP. DB Human immune/haematopoietic antigen PN W0200157182-A2. PD 04-A716-2001	PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 29.30% Best Local Similarity: 21.66% Query Match: 13.96%	RESULT 344 Percent Similarity; 47.00% Best Local Similarity; 26.00% Query Match: 13.96%	RESUL 545 Percent Similarity: 47.00% Best Local Similarity: 26.00% Query March: 13.96% PERTITY 546	AAD54480 standard Human CIP DNA #1. WO200299055-A2. 12-DEC-2002.	PA (EXEL-) EXELIXIS INC.  Percent Similarity: 40.22*  Best Local Similarity: 26.09*  Query Match: 13.96*	KESOLI 34)  ADL13850 standard; DNA; 180550 BP.  DE Osteoarthritis-associated polymorphic PN W02003054166-A2.  PD 03-IIII2003	SHE	ID AEQ75562 standard; DNA; 188888 BP. DE Human related CYP 27C1 clone RP11-30F3 PN WC200264765-A2	22-AUG-2002 (CYTO-) CYT rcent Similari st Local Simil st Local Simil sry Match:	ID AEK84349 standard; cDNA; 222930 BP. DE Human cDNA differentially expressed PN WO200228999-A2.	11-APR-2002 (GENE-) GEN rcent Similari st Local Simil ery Match:	32 standard; c EST associate 137139-A1. -2002.	PA (TACM) MAINIBLANGAN N. PA (TACM) TAC N. PA (WARR) WARREN W C. Percent Similarity: 46.48% Best Local Similarity: 28.17% Query Match: 13.85%	ABV12569 standard; cDNA; 416 BP. DE Human prostate expression marker PN W0200160860-A2.

```
N; 511 BP. copathogen resistance-related CDNA - SEQ ID 1104.
 u; 511 BP. opathogen resistance-related cDNA - SEQ ID 2028.
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 IA; 456 BP.
Imour protein, SEQ ID No 1205.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
DICTIVE MEDICINE INC.
05% Conservative:
87% Mismatches:
185% Indels:
 DICTIVE MEDICINE INC.
05% Conservative:
87% Mismatches:
185% Indels:
 DDICTIVE MEDICINE INC.
05% Conservative:
87% Mismatches:
185% Indels:
 DICTIVE MEDICINE INC.
05% Conservative:
87% Mismatches:
Indels:
 IA; 461 BP.
sion marker cDNA 33694.
 ion marker cDNA 42596.
 IA; 434 BP.
ion marker cDNA 3391.
 A; 625 BP.
SEQ ID NO:772.
 .; 774 BP.
equence #780.
 IOTECH INC.
 A; 618 BP.
 34
83
85
85
85
 9338
 75.4
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Drosophila melanogaster expressed polynucleotide SEQ ID NO 18260. 27-SEP-2001.
 Drosophila melanogaster genomic polynucleotide SEQ ID NO 14965. W0200171042-A2. 27-SEP-2001. (PEKE) PE CORP NY.
 protein, CryET53
 9
27
28
 27
 8
27
12
 AAA89220 standard; DNA; 3501 BP.
Bacillus thuringiensis delta-endotoxin gene 158C2b.
US6150589-A.
 human diagnostic protein #18933
 ABA02133 standard; cDNA; 3006 BP.
Human zinc ion transport protein 26-encoding cDNA.
WO200181539-A2.
 AAVI5221 standard; DNA; 3501 BP.
Bacillus thuringiensis 158C2b toxin encoding DNA.
Bacillus Thuringiensis 158C2b toxin encoding DNA.
B35723758-A.
G3-MAR-1998.
(MYCO) MYCOGEN CORP.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 B. thuringiensis DNA encoding a toxic crystal WO200119859-A2.
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Indels:
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 Indels:
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 Indels:
 PD WO200181539-Az.
PD 01-NOV-2001.
PD A (BIOW-) BIOWINDOW GENE DEV INC SHANGHAI.
PERCENT Similarity: 41.79$ Conser
Best Local Similarity: 29.85$ Mismat
I3.85$ Indele
 CDNA; 2918 BP
 ABX34815 standard; cDNA; 3666 BP.
Human mddt cDNA SEQ ID 376.
 ABL07926 standard; cDNA; 3991 BP.
 BP
 AAS02474 standard; DNA; 3507 BP
 T 569
ABL21164 standard; DNA; 3003
 PN WO200119859-AZ.
PD 22-MAR-2001.
PA (MONS) MONSANTO CO.
PECCENT Similarity: 45.45%
Best Local Similarity: 29.87%
 PN "C.Z. PD 10-0CT-2002.
PA (INCY-) INCYTE GENOMICS INC.
Percent Similarity: 49.43%
Best Local Similarity: 29.89%
 PD 19-FEB-2004.

PA (EXEL-) EXELIXIS INC.
Percent Similarity: 41.79%
Best Local Similarity: 29.85%
Query Match: 13.86%
RESULT 568
 33.33% 24.07%
 Percent Similarity: 42.11%
Best Local Similarity: 32.63%
Query Match: 13.85%
 13.85%
 Percent Similarity: 45.45%
Best Local Similarity: 29.87%
Query Match: 13.85%
 Percent Similarity: 45.45%
Best Local Similarity: 29.87%
 13.85%
 21-NOV-2000.
(MYCO) MYCOGEN CORP.
 DNA encoding novel WO200175067-A2.
 AAS83129 standard;
 PD 11-OCT-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity:
 Best Local Similarity:
 PN US5723758-A.
PD 03-MAR-1998.
PA (MYCO) MYCOGEN
Percent Similarity:
 WO200279449-A2.
 Query Match:
RESULT 569
 Query Match:
 Query Match:
RESULT 573
 Query Match:
RESULT 574
 Query Match:
RESULT 575
 Human prostate cancer antigen nucleotide sequence SEQ ID NO:249.
W0200055174-A1.
 ABL21165 standard; DNA; 1003 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 14968.
W020011042-A2.
27-SEP-2001.
(PEKE) PE CORP NY.
 8
14
14
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27
28
 13
21
35
 16
35
18
 AAA78540 standard; DNA; 2029 BP.
Plant SDF polynucleotide sequence SEQ List 1 NO:372.
WO200040695-A2.
 16
28
16
 14
23
16
 Query Match:
13.85* Indels: 35
RESULT 563
ID AACS3418 standard; DNA; 1266 BP.
DE Arabidopsis thaliana DNA fragment SEQ ID NO: 74455.
PN EP1033405-A2.
 AAC32883 standard; DNA; 1364 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 996.
EP1033405-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Mismatches:
Indels:
 Mismatches:
 Human prostate expression marker cDNA 28044
 PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Percent Similarity: 41.05% Conservat
Best Local Similarity: 27.37% Mismatche
Query Match: 13.85% Indels:
 ADL22897 standard; DNA; 2476 BP.
Human MP2153 coding sequence SEQ ID NO: 17.
WO2004015069-A2.
 Indels:
 Indels:
 Indels:
 Indels:
 ABQ67685 standard; DNA; 774 BP.
Listeria innocua DNA sequence #487.
WO200228891-A2.
 ABV28053 standard; cDNA; 1131 BP
 AAF15814 standard; cDNA; 2422 BP
 (INSP) INST PASTEUR.
(CNRS) CNRS CENT NAT RECH SCI.
 (CNRS) CNRS CENT NAT RECH SCI.
 21-SEP-2000.
(HUMA-) HUMAN GENOME SCI INC.
(ROSE/) ROSEN C A.
PD 11-...
PA (INSP) INS1 ...
PA (CNRS) CNRS CENT NA1 ...
Percent Similarity: 45.10$
Rest Local Similarity: 29.41$
13.85$
 ID ABQ67685 standard; DNA; 774
DE Listeria innocua DNA sequence
N W020022891-A2.
DD 11-APR-2002.
PA (INSP) INST PASTEUR.
PA (CNRS) CNRS CENT NAT RECH SPECCENT Similarity: 45.10%
Dest Local Similarity: 45.10%
Query Match: 13.85%
RESULT 561
 PN WOLUNDSONG.
PD 13-JUL-2000.
PD (CRRE-) CERES INC.
Percent Similarity: 46.58%
Best Local Similarity: 27.40%
 41.76%
24.18%
13.85%
 41.76%
24.18%
13.85%
 46.34%
 Sest Local Similarity:
 Percent Similarity:
Best Local Similarity:
 Local Similarity:
 ID AAC53418 standar
DE Arabidopsis thal
PN EP1033405-A2.
PD 06-SEP-2000.
Percent Similarity:
 ID AAR15814 standar
DE Human prostate of
PN W020055174-A1.
PP 41-SEP-2000.
PA (HVMA-) HVMAN GE
PA (ROSE/) ROSEN C
Percent Similarity:
 06-SEP-2000
 23-AUG-2001
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Query Match: RESULT 565

Query M. RESULT

Match:

Best

Query Match: RESULT 564

Query Match: RESULT 562

				1	0	r
PA (PEKE ) PE CORP NY. Percent Similarity: 52.50% Best Local Similarity: 40.00% Query Match: 13.85%	Conservative: Mismatches: Indels:	5 11 8	DE MOUSE FYN CATCINOMA ASSO PN WC2003057146-A2. PD 17-JUL-2003. PA (SAGR-) SAGRES DISCOVERY	ciated .	gene, sky ib NO:11	133.
RESULT 576  ID ABL18500 standard; DNA; 4063 BP. DE Drosophila melanogaster genomic polynuclen melanogaster genomic polynuclen	otide	SEQ ID NO 6973.	Percent Similarity: Best Local Similarity: Query Match:	47.37% 33.33% 13.85%	Conservative: Mismatches: Indels:	1 2 9
NY. 42.11% Y: 32.63% 13.85%	Conservative: Mismatches: Indels:	9 27 28	ID ALB 2043 Standard; DNA; 'DB Mouse Fyn gene. PN WO2003008583-A2. PD 30-JAN-2003. PA (SAGR-) SAGRES DISCOVERY	DNA; 73634 BP.		
rd; DNA; 4650 BP. olynucleotide SEQ	ID NO 10965.		Percent Similarity: Best Local Similarity: Query Match: RESULT 585 ID ADB95863 standard:	47.37% 33.33% 13.85% DNA: 73634 BP.	Conservative: Mismatches: Indels:	1 29
	Conservative: Mismatches: Indels:	8 17 18	yn gene 39484-A2 2003. SAGRES larity: imilarit	DMic DNA Second Covery. 47.37% 33.3%	ce. Conservative: Mismatches:	88 7 ,
Coccus polynucleotide SEQ 2002. CHIRON SPA.	ID NO 10237.				encoding cDNA	SEQ II
PA (GENO-) INST GENOMIC RES. GENO-) INST GENOMIC RES. Best Local Similarity: 36,76% QUETY March: 13.85% BESTIM 479	Conservative: Mismatches: Indels:	8 17 18	PD 23-MAY-2002. PA (UYA) UNIV YALE. Percent Similarity: Best Local Similarity:	42.178 28.928 13.858	Conservative: Mismatches: Indele:	11 35
525 standard; DNA; 4734 BP. tococcus polynucleotide SEQ 234771-A2.	ID NO 8963.		RESULT 587 ID AAX91990 standard; DE Nucleotide sequenc PN W09927105-A2.	DNA; 1230025 e of the compl	gen	lamyd:
PA (CHIR-) CHIRON SPA.  PA (GENO-) INST GENOMIC RES.  Percent Similarity: 48.53%  Best Local Similarity: 36.76%  Query Match: 13.85%	Conservative: Mismatches: Indels:	8 17 18	PD 03-UTN-1999. PA (GEST ) GENSET. Percent Similarity: Best Local Similarity: Query Match:	38.618 24.75% 13.85%	Conservative: Mismatches: Indels:	330 350 370
1 standard; DNA; 4770 BP. coccus polynucleotide SEQ 4771-A2.	ID NO 6295.		RESULT SHE PERCENT SIMILARITY: Best Local Similarity: Query Match:	48.53% 36.76% 13.85%	Conservative: Mismatches: Indels:	8 17 18
PD 02-PM1-2002. PA (CHIR-) CHIRON SPA. PA (GENO-) INST GENOMIC RES. Percent Similarity: 48.53% Best Local Similarity: 36.76%	Conservative: Mismatches:	8 17	RESULT 389 Percent Similarity: Best Local Similarity: Query Match: RESULT 590	45.10% 29.41% 13.85%	Conservative: Mismatches: Indels:	8 14 14
13.85% 154 standard; CDNA; 5674 nucleic acid-associated 054219-A2.	ın	18 CDNA SEQ ID NO:92.	ID ABQ69245 standard; DE Listeria innocua D PN W0200228991-A2. PD 11-APR-2002. PA (INSP ) INST PASTE	l; DNA; 3011208 B DNA sequence #68 EUR.	т. Т.	
003. INCYTE GENOMICS INC. arity: 45.71% milarity: 30.00%	Conservative: Mismatches: Indels:	11 4 4	PA (CNRS ) CNRS CENT NAT RECH SY Percent Similarity: 45.10% Best Local Similarity: 29.41% Query Match: 13.85% RESULT 591	NAT RECH SCI. 45.10% 29.41% 13.85%	Conservative: Mismatches: Indels:	8 14 14
875 BP. nomic p	22875 BP. genomic polynucleotide S	SBQ ID NO 37555.		570 B tion	related cDNA SEQ	e e
PA (PEKE) PE CORP NY.  Percent Similarity: 40.48%  Best Local Similarity: 25.00%  RESULT Match: 13.85%	Conservative: Mismatches: Indels:	13 32 18	(PROC ) PROCTER & ccent Similarity: st Local Similarity: sry Match:	AMBLE CO. 44.62% 33.85% 13.85%	Conservative: Mismatches: Indels:	7 26 10
ID ADA02615 standard; DNA; 73634 BP.		_	ID AAC00776 standard;	CDNA; 350 BP.		

```
1230025 BP.
he complete genome of Chlamydia pneumoniae.
 83698 BP. e1-like 4 encoding cDNA SEQ ID NO 9.
 172570 BP. ntiation related cDNA SEQ ID NO 114.
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1 29
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141
Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
Indels:
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Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 73634 BP.
NA sequence.
 3011208 BP.
puence #684.
 73634 BP
 , 350 BP.
 CH SCI.
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Query Match:
RESULT 601
 Query Match:
RESULT 606
 Query Match:
RESULT 607
 Query Match:
 JT 596
ADKS3732 standard, DNA, 587 BP.
Plant DNA sequence which confers altered metabolic characteristic #1115.
WO2003020936-A1.
 NO 16187
 NO: 2018
 Π
 86.
 SEO ID
 SEO
 AAS30336 standard; cDNA; 588 BP.
DNA encoding novel prostate gland antigen, Seq ID No
WO200155447-A1.
 10
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24
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7
 #2
 11
34
25
 11
39
10
 9
29
 Human phospholipase A2-like enzyme encoding cDNA W0200231162-A2.
 polynucleotide
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 antigen cDNA
 Human secreted protein 5' EST, SEQ ID NO: 774
 Mismatches:
 Human prostate expression marker cDNA 48201.
 BP.
SEQ ID NO:7955.
 PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Percent Similarity: 40.96% Conservat
Best Local Similarity: 27.71% Mismatche
Query Match: 13.74% Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 Drosophila melanogaster expressed 97-cep.
 Human reproductive system related WO200155320-A2.
 AAS80715 standard; cDNA; 795 BP
 AAL02017 standard; cDNA; 588 BP
 ID ADK53732 stanuard,
DE Plant DNA sequence which configure wood03020936-A1.
PD 13-MAR-2003.
PA (DOWC) DOW CHEM CO.
PA (DOWC) DOW AGROSCIENCES LLC Percent Similarity: 40.00%
Best Local Similarity: 40.00%
 13-MAR-2003.
(DOWC) DOW CHEM CO.
(DOWC) DOW AGROSCIENCES LLC.
 Human cDNA clone (3'-primer)
EP1074617-A2.
 PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 46.94*
Best Local Similarity: 32.65*
Query Match: 13.74*
 CDNA; 546
 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 N WOZVOZ-
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 41.67%
Best Local Similarity: 29.17%
 38.64%
27.27%
13.74%
 38.54%
27.08%
13.74%
 Percent Similarity: 46 94%
Best Local Similarity: 32.65%
Query Match:
DE Human secreted prote
PN EP1033401-A2.
PD 06-SEP-2000.
PA (GEST) GENSET.
Percent Similarity:
Best Local Similarity:
 T 599
ABL07235 standard;
 PD 18-APR-2002.
PA (FARB) BAYER AG.
Percent Similarity:
Best Local Similarity:
 23-AUG-200
 02-AUG-2001
 Query Match:
RESULT 596
 Query Match:
RESULT 595
 Query Match:
RESULT 597
 Query Match:
RESULT 600
 Query Match:
RESULT 593
 Query Match
RESULT 598
 Query M
RESULT
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Probe #10143 for gene expression analysis in human cervical cell sample. W0200157278-A2. 09-AUG-2001. (MOLE-) MOLECULAR DYNAMICS INC.
 Probe #14096 used to measure gene expression in human placenta sample. WO200157272-A2.
 ABA65239 standard; DNA; 1728 BP.
Human foetal liver single exon nucleic acid probe #13544.
WO200157277-A2.
 RESULT 608
ID ABA47354 standard; DNA; 1728 BP.
DE Human breast cell single exon nucleic acid probe #6049.
 13
18
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 13
18
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 20
33
 11
24
7
 14
43
8
DNA encoding novel human diagnostic protein #16519 WO200175067-A2.
 Bread wheat thiamine biosynthetic enzyme 1 cDNA #5.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 CO E I.
Conservative:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 gene
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Mismatches:
 AAC55207 standard; cDNA; 1163 BP.
Human secreted protein gene 18 SEQ ID NO:28
WO200047602-A1.
 Indels:
 Indels:
 ABL53444 standard; DNA; 1202 BP.
Glycerol-3-phosphate dehydrogenase (GPD)
KR2001011674-A.
 Indels:
 Indels:
 Mushroom GPD gene.
KR2001011673-A.
15-FEB-2001.
(ROKR-) ROK RURAL DEV ADMINISTRATION.
 (ROKR-) ROK RURAL DEV ADMINISTRATION.

(ROKR-) ROK RURAL DEV ADMINISTRATION.

(ROKR-) ADMINISTRATION.
 ADB46123 standard; cDNA; 1215 BP
 ВР
 AAI20210 standard; DNA; 1728 BP
 AAI45410 standard; DNA; 1728 BP
 (MOLE-) MOLECULAR DYNAMICS INC.

Int Similarity: 50.00%

Local Similarity: 42.59%

MARCh: 13.74%
 09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
 PA (DUPO) DU PONT DE NEMOURS & Percent Similarity: 45.26*
Percent Similarity: 24.21*
Query Match: 13.74*
RESULT 604
 17-AUG-2000.
(HUMA-) HUMAN GENOME SCI INC.
 DNA; 1202
 41.98%
25.93%
13.74%
 50.00%
42.59%
13.74%
 50.00%
42.59%
13.74%
 48.33%
30.00%
13.74%
 Percent Similarity: 41.98%
Best Local Similarity: 25.93%
Query Match: 13.74%
 Percent Similarity: 41.38%
Best Local Similarity: 25.29%
Query Match: 13.74%
DE DNA encoding novel h W0200175067-A2.
PD 11-0CT-2001.
PA (HYSE-) HYSEO INC.
Percent Similarity:
Best Local Similarity:
 ABL53352 standard;
 PA (ROKR-) ROK RURAL)
Percent Similarity:
Best Local Similarity:
Query Match:
 PA (MOLE-) MOLECULAR I
Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
 PN WO200157272-A2.
PD 09-AUG-2001.
PA (MOLE-) MOLECULA
Percent Similarity:
 PA (HUMA-) HUMAN G
Percent Similarity:
 US6512164-B1.
28-JAN-2003.
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16-JAN-2003. (CURA-) CURAGEN CORP. rcent Similarity: 50.00* st Local Similarity: 42.59* sty Match: 13.74*	ADH72315 standar Human gene of th WO2003102155-A2. 11-DEC-2003.	(CURA-) CURAGEN CORP. ccent Similarity: 50.00% st Local Similarity: 42.59% sry Match: 13.74% sULT 618	ID ADH72307 standard; DNA; 1800 BP.  DE Human gene of the invention NOV62b SEQ ID NO:1203.  PN WO2003102155-A2.  PD 11-DEC-2003.	CURAGEN CORP. illarity: 50.00% Similarity: 13.74%	ADH/2309 Standar Human gene of th WO2003102155-A2. 11-DEC-2003.	(CURA-) CURAGEN CORP.  (Crent Similarity: 50.00% st Local Similarity: 42.59% sry Match: 13.74% SULT 620	LD AMANOVEN SEGRETARY, 1908 BF.  DE Murine Oct4 enhancer and promoter.  PN W0200056932-A2.  PD 28-SBF-2000.  PN (1700E.) INTIV GEORGIA DES FOUND INC	ccent Similarity: 32.00% st Local Similarity: 21.33% sry Match: 13.74%	ID AAH98759 standard; cDNA; 1927 BP.  DE Human EST-derived coding sequence SEQ ID NO: 616.  PN W0200154477-A2.  PD 02-AUG-2001.	PA (HYSE-) HYSEQ INC.  Percent Similarity: 44.44\$ Conservative: 12  Best Local Similarity: 29.63\$ Mismatches: 28  Query Match: 13.74\$ Indels: 17  RESULT 622	ID AAH99768 standard; cDNA; 1927 BP. DB Human protein encoding cDNA sequence SEQ ID NO:603. PN WO200153455-A2. PD 26-JUL-2001.	(HYSE-) HYSEQ I ccent Similarity: st Local Similarity sry Match:	1D AAD19218 Standard; DNA; 1931 BP.  DB Human CG95 (or C870) lipase DNA.  PN WO200179446-A2.  PD 25-OTT-2001.	44.44% 29.63% 13.74%	DE Human soft tissue sarcoma-upregulated DNA - SEQ ID 993. PN W0200404838-A2. PD 10-UUN-2004.
PN WO200157271-A2.  PD 09-AUG-2001.  PA (MOLE-) MOLECULAR DYNAMICS INC.  Percent Similarity: 50.00% Mismatches: 16  Query Match: 13.74% Indels: 11	0 standard; DNA; 1728 BP. 10806 for gene expression analysis in human 7274-A2.	rce st	RESULT 610 ID AAK39397 standard, DNA, 1728 BP. DE Human bone marrow expressed single exon probe SEQ ID NO: 13954. PN W020015726-A2.	PD 09-AUG-2001.  PA (MOLE-) MOLECULAR DYNAMICS INC.  PA (MOLE-) MOLECULAR DYNAMICS INC.  Conservative: 4  Best Local Similarity: 42.59% Mismatches: 16  Query Match: 13.74% Indels: 11	KESULI 6LI DD AAK13656 standard; DNA; 1728 BP. DE Human brain expressed single exon probe SEQ ID NO: 13647. PN WO200157275-A2.	PD 09-AUG-2001.  AA (MOLE-) MOLECULAR DYNAMICS INC. Percent Similarity: 50.00% Conservative: 4 Best Local Similarity: 42.59% Mismatches: 16 Query Match: 13.74% Indels: 11	KESULT 6.1.  ID A8538985 standard; DNA; 1728 BP.  DE Human liver single exon probe, SEQ ID No 13975.  PN WO200157273-A2.	rce st	RESULT 613 ID AA105914 standard; DNA; 1728 BP. DE Probe #5905 used to measure gene expression in human breast sample. PN WO200157270-A2.	PD 09-AUG-2001. PA (MOLE-) MOLECULAR DYNAMICS INC. Percent Similarity: 50.00% Conservative: 4 Best Local Similarity: 42.59% Mismatches: 16 Query Match: 13.74% Indels: 11	RESULT 614  ID ABS13484 standard; DNA; 1728 BP.  DE Human genome-derived single exon probe ORF from lung SEQ ID No 13475.  PN WO200186003-A2.	PD 15-NOV-2001. PD TS-NOV-2001. PERCENT SIMILATILY: 50.00% Conservative: 4 Best Local Similarity: 42.59% Mismatches: 16 Query_Match: 13.74% Indels: 11	KESULT 615  ID ADD18217 standard; DNA; 1800 BP.  DE Human molecule (MOL) protein MOL2c DNA sequence.  NO2030330384-A2.	PD 16-JAN-2003.  A (CURA-) CURAGEN CORP. Percent Similarity: 50.00% Conservative: 4  Best Local Similarity: 42.59% Mismatches: 16  Query Match: 13.74% Indels: 11	RESULT 618 ID ADD18215 standard; DNA; 1800 BP. DE Human molecule (MOL) protein MOL2b DNA sequence. FN WO2003003984-A2.

us-09-989-293a-377.rng.spdi

RP. Conservative: 4 42.59% Mismatches: 16 13.74% Indels: 11	cDNA; 2476 BP. ; cDNA SEQ ID NO 146.	ICS INC. Conservative: 15 7.39\$ Mismatches: 31 3.74\$ Indels: 41	DNA; 2601 BP. e of a human secreted polypeptide.	BEECHAM CORP. BEECHAM PLC. 50.00% Conservative: 4 42.59% Mismatches: 16 13.74% Indels: 11	31 BP.	CORP. Conservative: 4 50.00% Mismatches: 16 Y: 42.59% Mismatches: 16	indels: 31 BP. ike molecule 4 (TLR-L4)	CORP. Conservative: 4 50.00% Conservative: 4 42.59% Mismatches: 16 13.74% Indels: 11	NA; 2631 BP. protein MOL2a DNA sequence.	CORP. Conservative: 4 50.00% Conservative: 4 Y: 42.59% Mismatches: 16 13.74% Indels: 11	, DNA, 2631 BP. invention NOV62a SEQ ID NO:1201.	0.00% Conservative: 4 2.59% Mismatches: 16 3.74% Indels: 11	DNA; 2646 BP. ae polynucleotide seqid 4832.	TICS CC	cDNA; 2651 BP. ster expressed polynucleotide SEQ ID
PA (CURA-) CURAGEN CORP Percent Similarity: 5: Best Local Similarity: 4: Query Match:		PD 23-MAY-2002. PA (INCY-) INCYTE GENOMICS PECCENT Similarity: 37.39 Best Local Similarity: 24.35 Query Match: 13.74	AAH78202 standard; Nucleotide sequence WO200166690-A2.	PA (SMIK) SMITHKLINE PA (SMIK) SMITHKLINE PA (SMIK) SMITHKLINE Percent Similarity; Best Local Similarity; Query Match:	3 standard; OL2 cDNA. 6339-A2.	24-JAN-2002. (CURA-) CURAGEN cent Similarity: st Local Similarit	anda like -A2.	il H C	KESULI 63.7  ID ADD18195 standard; DNA; 2631  DE Human molecule (MOL) protein  PN WO200303984-A2.	3 k k G	5 standard ene of the 02155-A2.	PD 11-DEC-2003. PA (CURA-) CURAGEN CORP. Percent Similarity: 50.00% Best Local Similarity: 42.59% Query Match: 13.74%	7 standard; lla pneumoni 36-81.	26-AUG-2003. (GENO-) GENOME THER crent Similarity: st Local Similarity: sty Match: SULT 640	ID ABL07234 standard; cDNA DE Drosophila melanogaster PN W0200171042-A2. PD 27-SEP-2001.
12 28 17		12 28 17	ng plants SeqID 1711.	6 117 133	1548.	20 34 55		15 31 41		7 29 11	2 .	4 16 11	A #3.	11 34 25	. 60
Conservative: 12 Mismatches: 28 Indels: 17	oncer SeqID 17.	Conservative: 12 Mismatches: 28 Indels: 17	expressing plants SeqID	Conservative: 6 Mismatches: 17 Indels: 13	gene #1	Conservative: 20 Mismatches: 34 Indels: 45	.0001570.	Conservative: 15 Mismatches: 31 Indels: 41	7.	Conservative: 7 Mismatches: 29 Indels: 11	e SEQ ID NO: 93	Conservative: 4 Mismatches: 16 Indels: 11	cDNA #	Conservative: 11 Mismatches: 34 Indels: 25	ID NO:120
::	SeqID		ressing plants SeqID	Conservative: 6 8 Mismatches: 1 14 Indels: 1	ene #1	e :. ⊆ ₩ 4.	AESULI 0.20 DE Human cDNA encoding clone TESTI10001570. PN EP1308459-A2. PD 07-MAY-2003.	107ECHNOLOGY. 37.39% Conservative: 1 24.35% Mismatches: 3 13.74% Indels: 4	1, 82, 82 standard; DNA; 2286 BP. Prokaryotic essential gene #17477. W020027188-A2.	7.71	ID NO: 93	4 L L	#	38.54\$ Conservative: 1 27.08\$ Mismatches: 3 13.74\$ Indels: 2	NO:120

SEQ ID NO 16184.

ila melanogaster genomic polynuc 1042-A2. 2001. PE CORP NY. Parity: 41.30% Conse imilarity: 27.17% Misma	SULT 649 SULT 649 SULT 649 ADB62683 standard; CDNA; 3072 BP. Human CDNA encoding clone MESAN20021220. EP1948459-A2. 07-MAY-2003. (HELL-) HELIX RES INST.	Percent Similarity: 50.00\$ Mismatches: 4  Best Local Similarity: 50.00\$ Mismatches: 16  Query Match: 13.74\$ Indels: 11  RESULT 650  ID ADOL0173 standard; DNA; 3138 BP.  DE Human NOVX polypeptide coding sequence SEQ ID NO: 195.	PA (CURA-) CURAGEN CORP.  Percent Similarity: 38.54 Conservative: 11  Best Local Similarity: 27.08\$ Mismatches: 34  Query Match: 13.74\$ Indels: 25  RESULT 651 D ABA94700 standard; CDNA; 3197 BP.	04490-A2. 1-2002. 1-2002. 1-2002. 1-2002. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-2003. 1-20	RESULT 652  ID ABZ74133 standard; DNA; 3200 BP.  DE Secreted protein gene 191 genomic fragment HLYGE16, SEQ ID NO:1280.  PN WQ200277013-A2.	it it	ABZ74132 standard; DNA; Secreted protein gene 19; WO200277013-A2.	Percent Similarity: 51.72\$ Conservative: 7 Percent Similarity: 39.66\$ Mismatches: 21 Query March: 13.74\$ Indels: 7 RESULT 654	ID ADA98667 standard; DNA; 3200 BP.  DE Human secreted protein-related DNA sequence #260.  PN W02003004623-A2.  PD 16-JAN-2003.  DA (HTMA) -) HTMAN GENONE SCT INC	ccent Similarity: 51.72 st Local Similarity: 39.66 sry Match: 13.74 SULT 655	ID ADA98668 standard; DNA; 3200 BP. DE Human secreted protein-related DNA sequence #261. PN W02003004623-A2. PD 16-JAN-2003. PA (HTMA-) HUMAN GENOME SCI INC.	ccent Similarity: 51.72% st Local Similarity: 39.66% ory Match: 13.74% SULT 656
erve atck ls:	Vold SEQ ID NO:1207.  Conservative: 4  Mismatches: 16  Indels: 11	DP. 14845. Conservative: 7 Mismatches: 21	igen genomic seque	Construction of Minatches: 20 Indels: 5 F	Conservative: 6 Mismatches: 20	CDNA SEQ ID	Conservative: 12 Mismatches: 28 Indels: 17	p. gulated DNA - SEQ ID 5557.	C. Conservative: 12 Mismatches: 28 Indels: 17	P. #3016.	NTRUM. S.	Conservative: 4 Mismatches: 16 Indels: 11
(PEKE ) PE CORP NY coent Similarity: st Local Similarity: sty Match: SULT 641 ADH72311 standard;	DE HUMAN SENG OF THE INVENTION NOVESCENG SEQ PN WC2003102155-A2.  PD 11-DEC-2003.  PA (CURA-) CURACEN CORP.  Percent Similarity: 50.00% Const Best Local Similarity: 42.59% Misma Query, Match: 13.74% Inde-	RESULT 642  RESULT 642  D AAH16115 standard; cDNA; 2739 BP.  DE Human cDNA sequence SEQ ID NO:14845  PN EP1074617-A2.  PD 07-FEB-2017.  PA (HELI-) HELIX RES INST.  Percent Similarity: 51.72% COMMON MATCH.  DOCT. MATCH.	0 standard; DNA; 2740 mmune/haematopoietic 7182-A2. HUWAN GENOME SCI INC	Best Local Similarity: 39.23% Conservations Local Similarity: 39.22% Mismatches Query Match: 13.74% Indels: RESULT 644  ID AAK73991 standard; DNA; 2740 BP. DE Human immune/haematopoietic antigen genomic PN W0200157182-A2.	PD 09-AUG-2001.  PA (HUMA-) HUMAN GENOME SCI INC.  Percent Similarity: 50.98%  Best Local Similarity: 39.22%  Ousty Match.	SULT 645 SULT 645 ACC72857 standard; cDNA; 2747 Human cancer related protein 6 WO2003025138-A2.	PD 27-MARK-2003. PA (EOSB-) EOS BIOTECHNOLOGY INC. Percent Similarity: 44.44% Best Local Similarity: 29.63% Query Match: 13.74%	KESULI 646 ID AD022737 standard; DNA; 2789 BP. DE Human soft tissue sarcoma-upregulated DNA PN W20040468938-A2. PD 10-JUM-2004.	PA (PROT-) PROTEIN DESIGN LABS INC Percent Similarity: 44.44* Best Local Similarity: 29.63* Query Match: 13.74*	ADF82460 standard; DNA; 2796 Bi Leukaemia-related DNA sequence WO2003039443-A2.	PA (DEKR-) DEUT KREBSFORSCHUNGSZENTRUM. PA (UYLU-) UNIV LUDWIG MAXIMILIANS. PA (HAFEA, HAFERLACH T. PA (SCHO/) SCHOCH C. PA (KRRN/) KRRN W.	rcent Similarity. st Local Similari sry Match:

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WO2003062376-A2.
 Percent Similarity
 31-JUL-2003
 07-MAR-2002
 17-JUL-2003
 (WEBS/) (KETC/) (DFRA/) (BEAS/) (
 Query Match:
RESULT 668
 Query Match:
 Query Match:
RESULT 666
 Query Match:
 Query Match:
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 (PLA2) protein.
 ADF76730 standard; cDNA; 3986 BP.

Novel human secreted and transmembrane protein cDNA SeqID 405.

WO2003072035-A2.
Human secreted protein-related DNA sequence - SEQ ID No 299.
WO200277188-A2.
03-OCT-2002.
 - SEQ ID No 300
 ABZ67689 standard; DNA; 3200 BP.
Human secreted protein encoding genomic DNA SEQ ID NO 1212.
WO200277186-A2.
 genomic DNA SEQ ID NO 1213
 Human soft tissue sarcoma-upregulated DNA - SEQ ID 7717.
 15
43
 11
34
25
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 727
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 21
 A2
 ABLG5195 standard; DNA; 3777 BP.
Lung cancer related gene sequence SEQ ID NO:3532.
WO200194629-A2.
 DNA; 3587 BP.
818 amino acid phospholipase
 Conservative:
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Mismatches:
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Mismatches:
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Mismatches:
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 Mismatches:
Indels:
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 Human secreted protein-related DNA sequence WO200271188-A2.
 Indels:
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 Indels:
 WO2004048938-A2.
10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
 ABZ67690 standard; DNA; 3200 BP. Human secreted protein encoding WO200277186-A2.
 ADE31412 standard; DNA; 4013 BP.
 ADQ24897 standard; DNA; 4003 BP
 N WCACCT-2002.
PD 03-OCT-2002.
PA (HUMA-) HUWAN GENOME SCI INC.
Percent Similarity: 51.72%
Best Local Similarity: 39.66%
 03-OCT-2002.
(HUMA-) HUMAN GENOME SCI INC.
 (HUMA-) HUMAN GENOME SCI INC.
 PA (HUMA-) HUMAN GENOME SCI INC
Percent Similarity: 51.72%
 N WCZVUZZOO3.
PD 24-JUL-2003.
PD 14-JUL-2003.
Percent Similarity: 38.54%
Best Local Similarity: 27.08%
 51.72%
39.66%
13.74%
 PA (HUMA-) HUMAN GENORM 5172%
Percent Similarity: 51.72%
Best Local Similarity: 39.66%
Best Local Similarity: 13.74%
 Percent Similarity: 51.72%
Best Local Similarity: 39.66%
Query Match: 13.74%
 PD 04-SEP-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 41.18*
Best Local Similarity: 23.53*
Query Match: 13.74*
 Percent Similarity: 41.18%
Best Local Similarity: 23.53%
Onerv Match: 13.74%
 Percent Similarity: 41.18% 3est Local Similarity: 23.53% Query Match: 13.74%
 13-DEC-2001.
(AVAL-) AVALON PHARM.
 ADB61644 standard;
Human DNA encoding
WO2003060132-A1.
 Best Local Similarity:
Query Match:
 Query Match:
RESULT 660
 Query Match:
RESULT 661
 Query Match:
RESULT 659
 Query Matc
RESULT 663
 RESULT 657
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Drosophila melanogaster expressed polynucleotide SEQ ID NO 4076.
WO200171042-A2
 ABL17390 standard; DNA; 11359 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 3643.
WG20011042-A2.
Z7-SEP-2001.
(PEXE) PE CORP NY.
 NO:1313
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 1 12
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 Conservative:
Mismatches:
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 Mismatches:
 ADA02795 standard; DNA; 52479 BP.
Mouse Tnfsfil carcinoma associated gene, SEQ
WO2003057146-A2.
 Human prostate expression marker cDNA 25463.
WO200160860-A2.
 Human prostate expression marker cDNA 25841 WO200160860-A2.
 PD 33-AUG-2001.
PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Percent Similarity: 41.18% Conservat
Best Local Similarity: 23.53% Mismatche
 23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
3nt Similarity: 41.18% Conservat
Local Similarity: 23.53% Mismatche
13.74% Indels:
 Indels:
 ABL03198 standard; cDNA; 11390 BP.
 RESULT 671
ID ADB72533 standard; DNA; 52479 BP.
 ABV25472 standard; cDNA; 4688 BP
 AAD35032 standard; DNA; 52354 BP
 Human transporter protein gene.
US2002028773-Al.
 CDNA; 4688
PA (INCY-) INCYTE GENOMICS INC
Percent Similarity: 41.18%
 41.18%
23.53%
13.74%
 58.978
38.468
13.748
 Percent Similarity: 41.30%
Best Local Similarity: 27.17%
Ouery Match: 13.74%
 PA (SAGR-) SAGRES DISCOVERY.
Percent Similarity: 58.97%
Best Local Similarity: 38.46%
 39.478
 Best Local Similarity: 24.56%
Query Match: 13.74%
 13.74%
 DE MOUSE This still gene.
PN WO2003008583-A2.
PD 30-JAN-2003.
PA (SAGRE) SAGRES DISCOVERY.
Percent Similarity: 58.97%
 (GUES/) GUEGLER K.
(WEBS/) WEBSTER M.
(KETC/) KETCHUM K A.
(DFRA/) DI FRANCESCO V.
(BRAS/) BEASLEY E M.
 PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 41
Best Local Similarity: 27
Query Match: 13
 Best Local Similarity:
Query Match:
RESULT 665
 ABV25850 standard;
 Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
```

H . O D . DE K	ERY978748748978978978978978748748748668668668748668668668668	ervative: 1s: 1s: 1c acid #3 acches: 1s: complete ervative: acches: 1s:	8 15 1 15 1 15 15 15
trch: intrch:	122186 BP 122186 BP 216215 Bi 216215 Bi 326014 BP 526014 BP 526014 BP 526014 BP 526014 BP	Mismatches: Indels: D coding sequence. Conservative: Mismatches: Indels: BP. Conservative: Mismatches: Indels: BP. Conservative: Mismatches: Indels: Mismatches: Indels: Mismatches: Indels:	16 27 27 3 3 16 11 10 25 21 25 21 25 21

```
REBULT 684

MARGA186 standard; CDNA; 586 BP.

DE Rat sequence differentially expressed in response to a hepatotoxin #1093.

PN WO200210453-A2.

PA (GENE-) GENE LOGIC INC.

Percent Similarity: 37.62$ Mismatches: 27

CHEST LOCAL Similarity: 26.73$ Mismatches: 36

RESULT 685

ID ADB57405 standard; DNA; 586 BP.

DE TOXICITY-related gene, SEQ ID 2431.

PN WO2003064624-A2.

PD 07-AUG-2003.

PA (GENE-) GENE LOGIC INC.

Percent Similarity: 26.73$ Mismatches: 27

Mismatches: 27

Mismatches: 27
 ID 6426
 11
32
18
 DESULT 686

RESULT 686

ID ANAVESBS standard; CDNA; 648 BP.

DE Human procestete expression marker CDNA 46579.

PN WO200160860-A2.

PN WO200160860-A2.

PD 23-AUG-2001.

PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Percent Similarity: 47.37% Conservative: 11

Best Local Similarity: 28.07% Mismatches: 26

Occ. Match: 13.64% Indels: 4
 \begin{array}{c} 11\\ 32\\ 18\\ \end{array}
 11
27
36
 11
26
 8
14
10
 ID AB29936 standard; DNA; 321 BP.

DE N. gonorrhoeae nucleotide sequence SEQ ID 4461.

PN W0200279243-A2.

PD 10-0CT-2002.

PA (CHIR-) CHIRON SPA.

Percent Similarity: 25.61$ Mismatches: 39.02$ Mismatches: 13.64$ Indels: 14.64$ Indels: 15.64$ Mismatches: 15.64
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Human Boft tissue sarcoma-upregulated DNA - SEQ WO2004048938-A2.
10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
 Query Match: 13.64%
RESULT 687
ID ADQ23606 standard; DNA; 680 BP.
 ADR02302 standard; DNA; 729 BP.
PA (CHIR-) CHIRON SPA.

Percent Similarity: 39.02%
Best Local Similarity: 25.61%
Query Match: 13.64%
 Percent Similarity: 51.02%
Best Local Similarity: 54.69%
Query Match: 13.64%
RESULT 688
 37.62%
26.73%
13.64%
```

```
ADI43922 standard; DNA; 1500 BP
PN W02003023009-A2.
PD 20-MRA-2003.
A (INCY-) INCYTE GENOMICS INC.
Percent Similarity: 38.20%
Best Local Similarity: 22.47%
 02-JAN-2003
 Best Local Sin
Query Match:
RESULT 697
 Query Match:
RESULT 702
 Query Match:
RESULT 699
 Query Match:
RESULT 700
 Query Match:
RESULT 701
 Query Match:
RESULT 703
 Query Match:
RESULT 704
 Query Match:
 P P P P
 B
 ABK68669 standard; DNA; 963 BP.
Human DNA for olfactory and pheromone G protein-coupled receptor #269.
WO200224726-A2.
 ACC49343 standard; cDNA; 1097 BP.
Human G protein-coupled receptor GCREC-17 encoding cDNA SEQ ID NO:39.
 SEQ ID NO:1418.
 sequence
 10
22
4
 14
23
32
 14
23
32
 14
23
32
 13
41
22
 RESULT 691
ID ADC27651 standard; DNA; 872 BP.
DE Human colon specific nucleic acid (CSNA) Seg ID20.
PN W02003020953-A2.
 ABSS8843 standard; DNA; 977 BP.
Human G-protein coupled receptor (GPCR) gene #97.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 ABN91955 standard; DNA; 948 BP.
Staphylococcus epidermidis ORF nucleic acid
US6380370-B1.
 ADG83531 standard; DNA; 963 BP.
Human Olfactory and pheromone GPCR DNA #123.
 Indels:
 Indels:
 Indels:
 Indels:
 DE Prog embryonic gene sequence 09924933.
PN US2002081610-A1.
PD 27-UTN-2002.
PA (UYRQ) UNIV ROCKEFELLER.
Percent Similarity: 37.62% Mism Dest Local Similarity: 24.75% Mism Query Match:
 PN USBELLOCAL SAMPLAND TO 29-MAY-2001.
PA (SYNG-) SYNGENTA PARTICIPATIONS AG. Percent Similarity: 46.67% C Best Local Similarity: 30.67% N 13.64% I
 A. gossypii genomic DNA PAG1610UP
US6239264-B1.
 30-APR-2002. (GENO-) GENOME THERAPEUTICS CORP.
 ВР
 CDNA; 771
 PN W0200224726-AZ.
PD 28-MAR-2002.
PA (CHEM.) CHEMCOM SA.
Percent Similarity: 38.20%
Best Local Similarity: 22.47%
 PN US2003221205-A1.
PD 27-NOV-2003.
PA (VETTY) VETTHEN A.
Percent Similarity: 38.20%
Best Local Similarity: 22.47%
 DNA; 753
 38.20%
22.47%
13.64%
 Best Local Similarity: 45.71%
Query Match: 13.64%
 50.00%
30.77%
13.64%
 P5-9 single chain Fv DNA.
WO200171005-A2.
27-SEP-2001.
 Percent Similarity: 44.168
Best Local Similarity: 28.578
 13.64%
 WO200259313-A2.
01-AUG-2002.
(CURA-) CURAGEN CORP.
 PD 13-MAR-2003.
PA (DIAD-) DIADEXUS INC.
Percent Similarity: 45
 PN WO20025913-A2.
PD 01-AUG-2002.
PA (CURA-) CURAGEN CORP
Percent Similarity: 3
Ouery Match:
RESULT 696
 AAS97141 standard;
 ABS76754 standard;
 Percent Similarity:
Best Local Similarity:
 (KUFE/) KUFER P.
 Percent Similarity:
 Query Match:
RESULT 693
 Query Match:
RESULT 694
 Match:
 689
 RESULT 690
 Query M
RESULT
 RESULT
 Query
 E E
```

```
AAD05663 standard; cDNA; 1500 BP.
Arbbidopsis thaliana ABA (abscisic acid)-insensitive 4, ABI4 cDNA.
WC200136586-A2.
25-MAY-2001.
 15
40
14
 15
40
14
 15
40
14
 16
31
26
 14
23
32
 14
23
32
 AACS4977 standard; DNA; 1362 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 79743.
EP1033405-A2.
 AAS69004 standard; cDNA; 1422 BP.
DNA encoding novel human diagnostic protein #4808.
WO200175067-A2.
 PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.

PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

Percent Similarity: 38.20% Conservative:

Best Local Similarity: 22.47% Mismatches:

Query Match:

13.64% Indels:
Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Plant yield-related polynucleotide clone G625 W02003013227-A2. 20-FEB-2003. (MEND-) MENDEL BIOTECHNOLOGY INC.
 Mismatches:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 AAZ37308 standard; DNA; 1499 BP. Arabidopsis Abi4 coding sequence WO9955840-Al.
 Arabidopsis Abi4 coding sequence W09955840-A1.
 PA (PION-) PIONEER HI-BRED INT INC.
Percent Similarity: 41.94*
Best Local Similarity: 25.81*
Query March: 13.64*
 ADD30509 standard; cDNA; 1500 BP
 JT 698
ADC86246 standard; DNA; 1363 BP.
Human GPCR gene SEQ ID NO:699.
EP1270724-A2.
 AAZ37307 standard; DNA; 1500 BP
 DE Arabidopsis Abi4 coding sequence by W09955840-Al.
PD 04-NOW-1999.
PA (REGC) UNIV CALIFORNIA.
PA (GEHO) GEN HOSPITAL CORP.
Percent Similarity: 25.81%
Best Local Similarity: 25.81%
 WOSPSSCSCS

WOOSPSSCSCSCS

(REGC) UNIV CALIFORNIA.

(GEHO) GEN HOSPITAL CORP.
 PA (VENUE)
Percent Similarity: 41.945
Best Local Similarity: 25.81$
13.64$
 40.66%
24.18%
13.64%
 40.00%
23.16%
13.64%
 41.94% 25.81%
 13.64%
 13.64%
 PD 11-OCT-2001.

PA (HYSE-) HYSEQ INC.

Percent Similarity:

Best Local Similarity:
 PD 06-SEP-2000.
Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
```

e #1550.	15 40 14	14 29 20	5 23 12	5 23 12	10 22 22	15 27 13	
related polynucleotide	Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:	ID NO:25. Conservative: Mismatches: Indels:	ID NO:206. Conservative: Mismatches: Indels:	013890. Conservative: Mismatches: Indel8:	SEQ ID NO: 3. Conservative: Mismatches: Indels:	SEQ ID NO:3770
actor	25.81% M: 13.64% II: cDNA; 1668 BP.		951 BP. ne SEQ I)	1851 BP. gene SEQ II	NA; 2317 BP. clone LIVER200 str. TECHNOLOGY. c. 80%	O O	NA; 3153 BP. gene sequence
DE Plant transcription for US2004019277-A1.  PD 29-JAN-2004.  SHERN) SHERMAN B K.  PA (JIAN) JIANG C.  PA (HEAR) HEARD J B.  PA (HAAK) HARRE J B.  PA (CREE) CREELMAN T J.  PA (CREE) RACLIFFE O.  PA (RATC) RATCLIFFE O.  PA (REUP) REUBER T L.  PA (REUP) REUPER T L.	.0		ADJ67311 standard; DNA; 18 Human ovarian specific ger wo2004013311-A2. 12-FEB-2004. (DIAD-) DIADEXUS INC. cent Similarity: 38.60% t Local Similarity: 29.82% ry Match:	Mul 707  AbJ67492 standard, DNA,  Human ovarian specific g W02004013311-A2.  12-FEB-2004.  (DIAD-) DIADEXUS INC.  Sent Similarity: 29.82.  Local Similarity: 29.82.	ID ADB62669 standard; cl DE Human CDNA encoding of PN EP1308459-A2. PD (7-MAY-2003. PA (HELL-) HELIX RES ING PA (REAG-) RES ASSOC BIG Percent Similarity: 38 Best Local Similarity: 25 Query Match: 12	KESSULT 709  ID ADK65983 standard; ID ADK65983 standard; ID Rice NADPH-oxidase or WO200409820-A1.  PD 29-UAN-2004.  PA (BADI ) BASF PLANT & Percent Similarity: Best Local Similarity: Dest Local Similarity:	433 standard; cancer related 194629-A2. C-2001.
D	Percent Best Loc Query Ma RESULT 7 ID AAI DE Hum	PD 26-PD 26-PA (HY Percent Best Loc Query Ma	ID ADD DE HUM PN WOM PD 12- PA (DI Percent Best Loc	KESULI 707 DE ADJ67492 DE HUMAN OV PN WO200401 PD 12-FEB-2 PA (DIA) Percent Simil Best Local Si Query Match:	NESCHIA DE HUM PN EP1 PD 07- PA (HE PA (HE PA (RE PETCENT BEST LOC	KESULT 709  DE ADK6598  DE ADK6598  DE ADC046  PN WO20046  PA (BADI )  Percent Simi  Best Local S  Query Match:	ID ABL DE Lun PN WO2 PD 13-

```
ID AAH54495 standard, DNA; 3504 BP.
DE S. epidermidis genomic polynucleotide sequence SEQ ID NO:3859.
PN WO200134809-A2.
PD 17-MAY-2001.
PA (GLAX) GLAXO GROUP LTD.
PRACCIS Similarity: 28.57$
Best Local Similarity: 28.57$
RESULT 716
ID ADM67075 standard, DNA; 4152 BP.
DE Murine adipocyte specific DNA SeqID 210.
PN WO20040161618-A2.
PD (S-FEB-2004.
PA (HMGE-) HMGENE INC.
 ADB33468 standard; DNA; 4861 BP.
Human novel lung related polypeptide DNA SEQ ID NO 395.
US2003054368-A1.
 37
 9
37
12
 9
37
12
 9
37
12
 10
22
22
 16
46
8
 7
24
12
 Human origin of replication complex protein 1 gene. US5614618-A.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 AAT62360 standard; cDNA; 3214 BP.
Human origin of replication complex ORC1 gene.
WO9640977-Al.
 Mismatches:
Indels:
 Mismatches:
 ID ADM67075 standard; DNA; 4152 BP.

DE Murine adipocyte specific DNA SeqID 210.

PN WO2004011618-A2.

PD 05-FEB-2004.

PA (HMGE-) HMGENE INC.

Percent Similarity: 41.94* Conservat Best Local Similarity: 24.73* Mismatch Query Match: 13.64* Indels:
 Indels:
 Indels:
 Indels:
 ADB63192 standard; cDNA; 3231 BP.
Human cDNA encoding clone SPLEN20111450.
EP1308459-A2.
 DE Human autoimmune disorder gene #34.

DE Human autoimmune disorder gene #34.

PN US200228617-A1.

PD 11-DEC-2003.

PA (UYVA-) UNIV VANDERBILT.

Percent Similarity: 42.35$

Query Match:

RESULT 712.
 AAS30131 standard; DNA; 4861 BP.
Human lung antigen genomic DNA #201.
WO200155303-A2.
02-AUG-2001.
 ECOLITY 713

IN PART 7287

END Human origin of replication completed by USS614618-A.

PN USS614618-A.

PN USS614618-A.

PN USS614618-A.

PA (REGC) UNIV CALIFORNIA.

PA (COLD-) COLD SPRING HARBOR LAB.

PETCENT SIMILATITY: 31.76*

QUETY MATCH: 13.64*

RESULT 714

ID ADB63192 standard; CDNA; 3231 BP.

DE Human CDNA encoding clone SPLEN20*

PN PR1308459-AS.

PR (REAS-) RES ASSOC BIOTECHNOLOGY.

PA (RELL-) HELIX RES INST.

PA (RELL-) RES ASSOC BIOTECHNOLOGY.

Percent Similarity: 38.89*

QUETY MATCH: 25.00*

QUETY MATCH: 715.
 CDNA; 3214 BP
 (REAS-) RES ASSOC BIOTECHNOLOGY.
 LAB.
 PN WOS6409//-n.
PD 19-DEC-1996.
PA (COLD-) COLD SPRING HARBOR LAI
PA (REGC) UNIV CALIFORNIA.
Percent Similarity: 42.35%
Best Local Similarity: 31.76%
Query Match: 13.64%
RESULT 713
ID AAT73287 standard; cDNA; 32?
""""" origin of replication
 PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 41.94%
Best Local Similarity: 30.65%
Query Match:
 42.35%
31.76%
13.64%
Percent Similarity:
Best Local Similarity:
Query Match:
RESULT 711
```

22

Query Match RESULT 719

```
ABX34940 standard; cDNA; 329 BP.
ABX34940 standard; cDNA; 329 BP.
Bovine EST associated with lactation/muscle/fat deposition #105.
26-SEP-2002.
(BYAT) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON) TAO N.
(WARR/) WARREN W C.
 Pseudomonas sp heavy metal transporter encoding DNA ORF04245.
DE1934720-A1.
25.JAN-2001.
(TIGR-) FIGR INST GENOMIC RES.
 ADB47539 standard; CDNA; 496 BP.
Human cDNA upregulated in dendritic cells SEQ ID NO 239.
US2003134283-A1.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
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Mismatches:
 Conservative:
 Conservative:
 Alloiococcus otitis antigenic protein encoding WO2003048304-A2.
 Mismatches:
Indels:
 Mismatches:
 (QUIA-) QUIAGEN GMBH.
(GREP) GES BIOTECHNOLOGISCHE FORSCHUNG MBH.
(DRET) DEF BIOTECHNOLOGISCHE FORSCHUNG MBH.
(MRD1-) MEDIZINISCHE HOCHSCHULE HANNOVER.
 AAZ21474 standard; cDNA; 499 BP.
Phaseolus lunatus DAD1 nucleotide sequence.
WO9947689-A2.
 ADI.17994 standard; DNA; 499 BP.
DNA encoding lima bean OB-related protein.
US2004067520-Al.
 Indels:
 Indels:
 Indels:
 Indels:
 ш
 AAD46127 standard; DNA; 163350 BP.
 8
 ADB06399 standard; DNA; 468 BP
 (DUPO) DU PONT DE NEMOURS &
 PN WCZCUCZ...
D12-JUN-2003.
PA (AMHP) WYETH HOLDINGS CORP.
Percent Similarity: 43.02%
Best Local Similarity: 26.74%
 Human tumour suppressor gene WO200268468-A2.
 T 730
AAF26426 standard; DNA; 474
 38.89%
25.00%
13.64%
 PA (Marcont Similarity: 49.12%
Best Local Similarity: 37.29%
 43.06%
27.78%
13.53%
 Percent Similarity: 36.26%
Best Local Similarity: 25.27%
Query Match: 13.53%
 Percent Similarity: 47.83%
Best Local Similarity: 30.43%
Query Match: 13.53%
 PA (PEKE) PE CORP NY.
Percent Similarity: 51.43
Best Local Similarity: 42.86
 13.64%
 DE .

 06-SEP-2002.
(PEKE) PE CORP NY.
 Percent Similarity:
Best Local Similarity:
Query Match:
 08-APR-2004.
(MANN/) MANN M B.
 PA (DUPO) DU PON'
Percent Similarity:
 Query Match:
RESULT 733
 Query Match:
 RESULT 727
 2 2
 Human immune/haematopoietic antigen genomic sequence SEQ ID NO:42364.
WO200157182-A2.
 JT 725
MENUGO948 standard; cDNA; 23407 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 15326.
WO200171042-A2.
27-SEP-2001.
 Drosophila melanogaster expressed polynucleotide SEQ ID NO 37850.
WO200171042-A2.
27-SEP-2001.
 Human digestive system antigen genomic sequence SEQ ID NO: 3236.
WO200155314-A2.
 ABL20832 standard; DNA; 8298 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 13369.
27-SEP-2001.
 ABL20834 standard; DNA; 8673 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 13975.
WO200111042-A2.
 NO 13963
 Drosophila melanogaster genomic polynucleotide SEQ ID NO 4633.
WO200171042-A2.
27-SEP-2001.
 SEQ ID
 7
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
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Mismatches:
 ABL20830 standard; DNA; 12319 BP.
Drosophila melanogaster genomic polynucleotide
WO200171042-A2.
 Mismatches:
Indels:
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 Indels:
 Indels:
 ABL14456 standard; cDNA; 28918 BP
 ABL17720 standard; DNA; 12319 BP
 AAK87552 standard; DNA; 18188 BP
 AAK89660 standard; DNA; 6784 BP
 PN WO200155314-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 40.51%
Best Local Similarity: 27.85%
PD 20-MAR-2003.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 41.94%
Best Local Similarity: 30.65%
 PN WOLVELD PD 09-AUG-2001.
PD 09-AUG-2001.
PA (HUMAN GENOME SCI INC. Percent Similarity: 42.86% Pest Local Similarity: 28.57% Rest Local Similarity: 13.64%
 09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 ID ABL20032 standard; DNA; 8296
DE Drosophila melanogaster genc
PN WO20171042-A2.
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 40.58%
Best Local Similarity: 30.43%
Query Match: 13.64%
RESULT 721
 PD 27-SEP-2001.

PA (PEKE) PE CORP NY.

Percent Similarity: 40.58*

Best Local Similarity: 30.43*

13.64*
 PA (PEKE,). Percent Similarity: 43.00%
Rest Local Similarity: 28.00%
 Percent Similarity: 40.58% Best Local Similarity: 30.43% Query Match: 13.64%
 40.58%
 Best Local Similarity: 30.43%
 13.64%
 PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity:
 (PEKE) PE CORP NY
 Query Match:
RESULT 724
```

Query Matc RESULT 723

RESULT

10 33

12 27 9

0.8

NO:339

SEQ ID

DNA

13.53%

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Query Match:
RESULT 742
 Query Match:
RESULT 744
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:329. WO2003048304-A2.
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:327.02003048304-A2.
12-JUN-2003.
(AMMP) WYETH HOLDINGS CORP.
 ADB06391 standard; DNA; 855 BP.
Aloiococcus otitis antigenic protein encoding DNA SEQ ID NO:331.
WC2003048304-A2.
12-JUN-2003.
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:335 WO2003048304-A2.
 27 6
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Query Match:
13.53$ Indels:
RESULT 734

ID ACHTASAS standard; DNA, 584 BP.
DE Human genome derived single exon probe #5631.
PN US2003194704-A1.
 DE HUMAN COLON CANCER CELL EXPRESSED CONA #208B.
PN US2002155438-Al.
PD 24-OCT-2002.
PA (SIMP) SIMPSON A J G.
PA (NETO/) NETO E D.
PA (BREN/) BRENTANI R R.
PA (BREN/) BRENTANI R R.
PA (SIMILARILY: 43.04% CONSERVATIV
BEST LOCAL SIMILARILY: 30.38% Mismatches:
Ouery Match: I3.53% Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 Fusarium venenatum EST SEQ ID NO:1476.
WO200056762-A2.
28-SEP-2000
 (NOVO) NOVO NORDISK BIOTECH INC (NOVO) NOVO NORDISK AS.
 AAF08953 standard; cDNA; 633 BP
 ACD93676 standard; cDNA; 757 BP
 ID ADBO6391 standard; DNA; 855 BP.
DE Alloicoccus otitis antigenic property 12-JUN-2003.
PA (AMHP) WYETH HOLDINGS CORP.
Percent Similarity: 43.02%
Best Local Similarity: 26.74%
 ADB06389 standard; DNA; 855 BP
 ADB06395 standard; DNA; 855 BP
 ID ACH72436 standard; www, creep Human genome derived single e PN US2003194704-A1.
PN US2003194704-A1.
PA (PENN/) PENN S G.
PA (RANK/) RANK D R.
PA (HANZ/) HANZEL D K.
PETCENT Similarity: 52.17%
Best Local Similarity: 41.30%
 PA (AMHP) WYETH HOLDINGS CORP.
Percent Similarity: 43.02%
Best Local Similarity: 26.74%
 PD 12-JUN 2003.
PA (AMHP) WYETH HOLDINGS CORP.
Percent Similarity: 43.02%
Best Local Similarity: 26.74%
Query Match: 13.53%
 RESULT 738
ID ADB06387 standard; DNA; 855
 40.32%
24.19%
13.53%
 43.02%
26.74%
13.53%
 PA (AMHP) WYETH HOLDLING
Percent Similarity: 43.02%
Best Local Similarity: 26.74%
13.53%
(HECH/) HECHT R I.
(PELL/) PELLEYMOUNTER M A.
(TOOM/) TOOMBS C F.
 int Similaricy:
Local Similarity: 30.43%
13.53%
 47.83$
 ID
DE FUS.
PN W0200.
PN W0200.
PD 28-SEP-20.
PA (NOVO) NOVC
P
PA (HECH/) HECHT R
PA (PELL/) PELLEYMO
PA (TOOM/) TOOMBS OF
PERCENT SIMILARITY;
Best Local Similarity
 12-JUN-2003
 Query Match:
RESULT 737
 Query Match:
```

```
Query Match:
13.53* Indels:
21
RESULT 747
ID AAZS255 standard; cDNA; 1458 BP.
DE Human secreted protein clone ykl43_1 nucleotide sequence SEQ ID NO:201.
PN W09958642-A2.
 Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:333. WO2003048304-A2.
 Drosophila melanogaster expressed polynucleotide SEQ ID NO 3161.
WO200171042-A2.
 1D ADB69278 standard; DNA; 984 BP.

DE C. neoformans coding sequence with introns SEQ ID NO:1044.

PN WOO00305205-A2.

PD 26-UNN-2003.

PA (ELIT-) ELITRA PHARM INC.

Percent Similarity: 37.50% Mismatches: 29

Query Match: 13.53% Indels: 21

RESULT 743
 13
41
7
 13
33
 14
34
15
 9
18
21
 4
10
11
 ABZ40187 standard, DNA; 1215 BP.

ABZ40187 standard; DNA; 1215 BP.

ABZ40187 standard; DNA; 1215 BP.

N. GONOFTHOGAE nucleotide sequence SEQ ID 4963.

PN WO200279243-A2.

PD 10-CCT-2002.

PA (CHIRA) CHIRN SPA.

Percent Similarity: 50.98%

Query Match: 13.53%

RESULT 746

ID ABL02893 standard

PN WO200171*

PN WO200171*

PN WO200171*
 8
24
 24
 Chlamydia trachomatis DNA sequence SEQ ID NO:414.
WO200208267-A2.
 Conservative:
Mismatches:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Mismatches:
Indels:
 Mismatches:
 Indels:
 ABS63259 standard; cDNA; 1140 BP.
High level promoter polynucleotide #6.
WO200261098-A2.
 Ξ.
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ც
 DNA; 1191 BP
 DNA; 1599 BP
ADB06393 standard; DNA; 855 BP
 PD 18-NOV-1999.
PA (GEMY) GENETICS INST INC.
Percent Similarity: 40.57%
Best Local Similarity: 28.30%
Query Match: 13.53%
RESULT 748
 PN MACK-CONTROLL OB NEMOURS PA (DUPO) DU PONT DE NEMOURS Percent Similarity: 42.17% Best Local Similarity: 13.53%
 41.79%
28.36%
13.53%
 PD 31-JAN-2002.
PA (CORI-) CORIXA CORP.
Percent Similarity: 47.50%
Best Local Similarity: 37.50%
Query Match: 13.53%
 PD 27-SEP-2001.

PA (PEKE) PE CORP NY.

Percent Similarity:

Best Local Similarity:
 ABL92621 standard;
 ABZ40196 standard;
```

Mismatches: Indels:

10 32 25

Conservative:

Mismatches:

Indels:

```
AAL03406 standard; DNA; 7809 BP.
Human reproductive system related antigen DNA SEQ ID NO: 6094.
WO200155320-A2.
 DE Human reproductive system related antigen DNA SEQ ID NO: 8621.

PD 02-AUG-2001.

PA (HUMA-) HUMAH GENOME SCI INC.

Percent Similarity: 30.12$ Mismacches: 27

Query Match: 13.53$ Indels: 19
 Human nervous system related polynucleotide SEQ ID NO 8055. WO200159063-A2.
 Human reproductive system related antigen DNA SEQ ID WO200155320-A2.
 DE Human testicular antigen encoding DNA fragment PN WO200155317-A2.
 ABL98498 standard; DNA; 15517 BP.
Human testicular antigen encoding DNA fragment
WO200155317-A2.
 ADN95876 standard; DNA; 5449 BP.
Human BEC/LEC-related gene sequence SeqID800.
WO2003080640-Al.
 AAN70608 standard, cDNA, 9202 BP. VISNA sheep lentivirus genome. FR2586427-A.
 Query Match:
RESULT 763
ID AALO5934 standard; DNA; 15517 BP.
 DNA; 15514 BP
 DE Human reproductive bybecom properties of MO200155320-A2.

PD 02-AuG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 40.32%

Best Local Similarity: 30.65%
 PN NCCCT-2003.
PN (LUDW-) LUDWIG INST CANCER RES.
PA (LICN) LICENTIA LTD.
Percent Similarity: 40.62%
Best Local Similarity: 30.21%
 DE Human testicular antigen encoci
PN WO20015517-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 44.58%
 (HUMA-) HUMAN GENOME SCI INC.
nt Similarity: 44.58%
Local Similarity: 30.12%
Match:
 16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 (HUMA-) HUMAN GENOME SCI INC.
 7809
 ID ABA15724 standard; DNA; 7809
DB Human nervous system related
PN WO200159063-A2.
PD 16-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC
Percent Similarity: 40.32%
Best Local Similarity: 30.65%
Query Match:
 PN FACELLY PACTEUR.
PA (INSP.) INST PASTEUR.
Percent Similarity: 50.00%
Best Local Similarity: 13.81%
 44.58%
30.12%
13.53%
28.36%
13.53%
 RESULT 759
ID ABA15724 standard; DNA;
 PN WOZO015520-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME
Percent Similarity: 4
Best Local Similarity: 3
Query Match: 1
RESULT 764
 AAL05933 standard;
 Best Local Similarity:
 Local Similarity:
 ID ABL98497 standar
DE Human testicular
PN W0200155317-A2.
PD 02-AUG-201.
PA (HUMA-) HUMAN GE
Percent Similarity:
 Query Match:
RESULT 757
 RESULT 760
 RESULT 758
 SEQ ID NO 3158.
 NO 29137.
 SEQ ID
 9
18
21
 9
26
15
 36
 30
 14
27
16
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29
21
 10
29
21
 Conservative:
Mismatches:
Indels:
 Drosophila melanogaster expressed polynucleotide WO200171042-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
 C. neoformans genomic DNA sequence SEQ ID NO:44.
WO2003052076-A2.
26-JUN-2003.
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 ADB68954 standard; DNA; 3147 BP.
C. neoformans genomic DNA sequence SEQ ID NO:81
WO2003052076-A2.
 ID ABLZ5888 standard; DNA; 4282 BP.
DE Drosophila melanogaster genomic polynucleotide
PN W0200171042-A2.
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 41.79% Conservative:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Mismatches:
 Indels:
 Indels:
 Indels:
 Indels:
 AAT12999 standard; cDNA to mRNA; 2982 BP.
 ADB62997 standard; cDNA; 2612 BP.
Human cDNA encoding clone PROST20082430.
EP1308459-A2.
 ADA52961 standard; cDNA; 2279 BP.
Human coding sequence, SEQ ID 529.
BP1293569-A2.
19-MAR-2003.
 Nitrate reductase coding sequence. JP08023978-A.
 PA (TOYW) TOYOTA CHUO KENKYUSHO KK.

PA (TOYT) TOYOTA JIDOSHA KK.

Percent Similarity: 46.25$

Best Local Similarity: 28.75$

Query Match: 13.53$
 Human bone remodelling gene #101.08426186-B1.
 INST.
BIOTECHNOLOGY.
 ABL02892 standard; cDNA; 3563 BP
 (HELI-) HELIX RES INST. (REAS-) RES ASSOC BIOTECHNOLOGY.
 ADB68917 standard; DNA; 2984 BP.
 PN BLL.
PD 19-MAR-ZULL
PA (HELL) HELLIX RES
PA (REAS-) RES ASSOC BIOTECHN:
Percent Similarity: 45.33$
Perc Local Similarity: 33.33$
TOTAL:
 PD 30-JUL-2002.

PA (INCY-) INCYTE GENOMICS INC.
PECCENT Similarity: 38.16%
Best Local Similarity: 26.32%
Query Match: 13.53%
 ID ADB62997 standard; cDNA, 261
DE Human cDNA encoding clone PR
DE D1308459-A2.
DD 07-MAY-2003.
PA (HELL) HELIX RES INST.
PA (REAS-) RES ASSOC BIOTECHNOI
Percent Similarity: 38.82%
Best Local Similarity: 32.94%
 N WCLOUS.

PD 26-JUN-2003.

PD 26-JUN-2003.

Percent Similarity: 37.50%

Best Local Similarity: 25.00%
 Percent Similarity: 41.79%
Best Local Similarity: 28.36%
Ouery Match: 13.53%
 PA (ELIT-) ELITRA PHARM INC.
Percent Similarity: 37.50%
Best Local Similarity: 25.00%
Ouery Match: 13.50%
 13.53%
 27-SEP-2001.
(PEKE) PE CORP NY.
 30-JAN-1996.
 Query Match:
RESULT 753
 Query Match:
RESULT 752
 Query Match:
RESULT 755
 Query Match:
RESULT 756
```

Query Matc RESULT 754

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6 13

Conservative: Mismatches: Indels:

Conservative:

Mismatches:

Indels:

Conservative: Mismatches:

RESULT 751

Indels:

NO: 3149

SEQ ID

NO: 8622

12 27 19

Conservative: Mismatches:

Indels:

NO: 3150.

11

SEQ

Conservative:

Mismatches: Indels:

Conservative:

(GENO-) GENOME THERAPEUTICS (AMHP) WYETH.  Freent Similarity: 52.17%  STUT 773  ADB98063 standard; DNA, 3376  HBM-related clone contig b52  WOO0092000-A2.	rce st st sty sury	PD 21-NOV-2002.  PA (GENO-) GENOME THERAPEUTICS CORP.  PA (AMHP) WYETH.  Percent Similarity: 52.17% Conservative: 5 Best Local Similarity: 41.30% Mismatches: 11 Query Match: 13.53% Indels: 11 RESULT 775 ID ABX13172 standard; DNA; 65464 BP.  DB Human gene encoding a Noelin-1-like secreted protein.  PN US2002173459-Al.	St.	TableMa associated made-like DNA 101 cancer 8268-82.  8101NVBNT INT AB. Conservative: 10 arity: 35.06\$ Mismatches: 27 milarity: 35.38\$ Indels: 14		RESULT 779 RESULT 779 RESULT 779 RESULT 780	RESULT 784  RESULT 784  ADB12064 standard; DNA; 1754382 BP.  DE Alloiococcus otitis entire genome sequence SEQ ID NO:6651.  PN WO2003048304-A2.  PN WO2003048304-A2.  PA (AMHP) WYETH HOLDINGS CORP.  Percent Similarity: 43.02\$ Mismatches: 34  Query Match: 13.53\$ Indels: 15  RESULT 782  Percent Similarity: 48.15\$ Conservative: 7  Best Local Similarity: 35.19\$ Mismatches: 9  RESULT 783  RESULT 783  RESULT 783
Imilarity: 30.12% 13.53% 13.53% 15.54% 15.518 BP. 15.20-A2. 15.001. 17.001. 18.11y: 44.58% 19.12% 19.12%	13.53* Indels: 9 standard; DNA; 15518 BP. esticular antigen encoding DNA fragment SE 5317-A2. HUMAN GENOME SCI INC. Conservative: 1arity: 30.12* Mismatches:	13.53% Indels: ; DNA, 21833 BP. etal system related polynucleoti DME SCI INC. 44.44% Mismatches: 13.53% Indels:	8 standard; coding novel 47140-Al. 2002. ROSEN C A. RUBEN S M.	Coent Similarity:  t Local Similarity:  t Local Similarity:  ty Match:  ADJ29648 standard  Human musculoskel  US200400488-Al.	15.JAN-2004.  (HUMA-) HUMAN GENOME SCI INC. (cent Similarity: 44.44% it Local Similarity: 31.11% sry Match: SULT 770 ABR82622 standard; DNA, 33769 BE	Human Hbw W02001773 18-0CT-20 (GENO-) ccent Simila st Local Sim sty Match:	DE Human high bone mass (HBM) polynucleotide clone #4.  DB Human high bone mass (HBM) polynucleotide clone #4.  PN W0200192891-A2.  PN W0200192891-A2.  PD 06-DEC-2001.  PA (GENO-) GENOME THERAPEUTICS CORP.  PA (UYCR-) UNIV CREIGHTON SCHOOL MEDICINE.  Percent Similarity: 52.17% Conservative: 5  Best Local Similarity: 41.30% Mismatches: 11  Query Match: 13.53% Indels: 11  RESULT 772  ID ACC45363 standard; DNA; 33769 BP.  DE Human HBM gene fragment #4.  PN W020022764-A2.

DNA for cancer treatment.

			. 0								. 99:		
13 30 30	#345.	10 37 9	NO:59	10 21 4		12 23 16		3 13		6 19	eq ID 4	111 24 20	11 24 20
Conservative: Mismatches: Indels:	P. rphic nucleotide	Conservative: Mismatches: Indels:	de (cdps) SEQ ID	Conservative: Mismatches: Indels:	tor segid 1453.	Conservative: Mismatches: Indels:	.069	Conservative: Mismatches: Indels:	ID NO:7578.	Conservative: Mismatches: Indels:	eted protein, S	Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:
40.00% /: 27.00% 13.53%	cd; DNA; 179651 BP. Issociated polymorphic	GENOMICS INC. 40.26% ty: 27.27% 13.53%	rd, cDNA, 280 BP.  ved polynucleotide	R V. B K. 53.70\$ 13.42\$	Standard, DNA; 294 BP. eeboox transcription factor 04. ONNER T W.	43.48% 7: 26.09% 13.42%	; cDNA; 425 BP. ide SEQ ID NO 3	IC. 43.75% ': 37.50% ': 13.42%	:d; cDNA; 595 BP.	SS INST. 44.78\$ 7: 35.82\$ 13.42\$	DNA; 889 BP. a novel secr	GENOME SCI INC. 44.30% .ty: 30.38% .ty: 13.42%	d, DNA; 889 BP. /nucleotide #456. A. M. S C. 3 C. 13.42%
Percent Similarity: Best Local Similarity Query Match:	ADL13813 standard; DNA; I: Osteoarthritis-associated WO2003054166-A2:	PD 03-JUL-2003.  PA (INCY-) INCYTE G Percent Similarity: Best Local Similarity Query Match:	ABL/1216 standard; c Corn tassel-derived US2001051335-Al.	PA (LALGY) LALGUDI PA (LALGY) LALGUDI PA (TYOL/) LY L PA (SHER/) SHERMAN Percent Similarity: Query Match: RESULT 786	ADQUAGO Standar Maize homeobox t US200412339-A1. 24-JUN-2004. (CONN/) CONNER T (HECK/) HECK G R	Percent Similarity: Best Local Similarity Query Match:	AAI83630 standard Human polynucleot WO200164835-A2.	PD 07-SEP-2001. PA (HYSE-) HYSEQ INC PErcent Similarity: Best Local Similarity: Ouery Match:	/88 AH10743 uman cDN P1074617	(HEI (HEI int S Loca Mat	AAS26287 standard; cl Human cDNA encoding of WO200155322-A2.	PD 02-AUG-2001. PA (HUMA-) HUMAN GER Percent Similarity: Best Local Similarity Query Match:	ABX73628 standard; DNA; 8: DE Human novel polynucleotid PN US2002132753-A1. PD 19-SEP-2002. PA (ROSE) ROSEN C A. PA (RUBE) RUBEN S M. PA (BARA/) BARASH S C. PA (BARA/) BARASH S C. Best Local Similarity: 30.38% Query Match: 13.42%
Per Bes Que	SED	PD Per Bes	NES PN	PA PA Per Des RES	PA PA	Perce Best Query	2088	PD 0 PA ( Percen Best L Query	PN	PA PA Perce Best Query	S E E E	PD PA Perce Best Query	AESULIA

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AAK79093 standard; DNA; 988 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:33905.
WO200157182-A2.
 DE ADQ14965 standard; DNA; 1082 BP.
DE Oligonuclectide for detecting cytosine methylation SEQ ID NO 1556.
DE NO 001086312-A2.
DA (SPIG-) EPIGENOMICS AG.
Percent Similarity: 56.10* Conservative: 9
Best Local Similarity: 34.15* Mismatches: 10
Query_Match: B
 Oligonucleotide for detecting cytosine methylation SEQ ID NO 1555. WO200218632-A2.
 8
25
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 9
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121
 6
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38
17
 13
38
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 ID AD067219 standard; DNA; 2197 BP.

DE CORN AD1-like (ZM-MADS PRO1) promoter sequence.

PN W0203078590-A2.

PD 25-SEP-2003.

PA (PION-) PIONEER HI-BRED INT INC.

Percent Similarity: 41.05% Conservative:

Best Local Similarity: 27.37% Indels:

RESULT 79%

ID AAT08552 standard; CDNA; 2336 BP.

DE Oncogene R-zes mutant CDNA (exon 1, intron A).

PN W0953223-A1.
 Conservative:
Mismatches:
Indels:
 Conservative:
 ADD67219 standard; DNA; 2197 BP.
Corn Ap1-like (ZM-MADS PRO1) promoter sequence.
WO2003078590-A2.
 Mismatches:
Indels:
 DE Extented sequence for mouse IMX5_49.

DE Extented sequence for mouse IMX5_49.

PN W0200231116_A2.

PD 18_APR-2002.

PA (DIGI-) DIGITAL GENE TECHNOLOGIES INC.

Percent Similarity: 46.25% Conservation of the conservation of t
 ID AAT08552 standard, C. DE Oncogene R-ras mutant cDNA (exon 1, 110120-PN W0953223-A1.
PN W0953223-A1.
PN (USSH) US DEPT HEALTH & HUMAN SERVICES.
PA (USSH) US DEPT HEALTH & HUMAN SERVICES.
Percent Similarity: 39.33 Conser Best Local Similarity: 29.21 Mismat Indele
 ID AAH1992 standard; CDNA; 1922 BP.
DE Human CDNA sequence SEQ ID NO:14495.
PN EP1074617-A2.
PD 07-PEB-2001.
PA (HELL:) HELIX RES INST.
Percent Similarity: 44.78% Conse Best Local Similarity: 13.42% Misma Query Match:
 SULT 795
ADE61223 standard; DNA; 1789 BP.
Human gene U35246, SEQ ID NO 7141.
W02003016475-A2.
27-FEB-2003.
(GEHO) GEN HOSPITAL CORP.
(FARB) BAYER AG.
ID AAK79093 standard, L.v., ---
DE Human immune/haematopoietic antiç
PN W0200157182-A2.
PD 09-A0G-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 32.86%
Best Local Similarity: 13.42%
 ABQ14964 standard; DNA; 1082 BP.
 ID ADE61223 standard,,
DB Human gene U35246, SEQ ID NO
PN W02003016475-A2.
PD 27-FEB-2003.
PA (GEHO) GEN HOSPITAL CORP.
PA (FARB) BAYER AG.
PErcent Similarity: 38.37%
Best Local Similarity: 27.91%
 D ABO14964 standard, DAG, LUCK, DE Oligonucleotide for detecting DE Oligonucleotide for detecting PN WO200218632-A2.
PA (EPIG-) EPIGENOMICS AG. Percent Similarity: 56.10% Best Local Similarity: 34.15%
 Query Match:
RESULT 793
 Query Match:
RESULT 794
 Query Match:
RESULT 792
 Query Match:
RESULT 795
 Query Match:
RESULT 796
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ABLO6694 standard; cDNA; 5069 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 14564.
 DE Drosophila melanogaster expressed polynucleotide SEQ ID NO 14510.
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 38.27% Conservative: 8
Best Local Similarity: 28.40% Mismatches: 30
 Pneumocystis carinii major surface glycoprotein gene HMSG14. WO200009760-A1.
 Human cDNA differentially expressed in lung cancer #377.
US2003065157-A1.
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 33
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33
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22
 Aures427 standard; DNA; 3591 BP.
Human breast-specific protein coding sequence #26.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 AUFU7683 standard; DNA; 4355 BP.
Human secreted protein encoding DNA, seq id 166.
 Conservative:
Mismatches:
 Conservative:
Mismatches:
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Mismatches:
 Conservative:
Mismatches:
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 Conservative:
Mismatches:
 Mismatches:
DE Pneumocystis carini major surface glycopr
PN W0200009760-A1.
PD 24-FEB-2000.
PA (USSH) US DEPT HEALTH & HUMAN SERVICES.
Percent Similarity: 50.60% Conserval
Best Local Similarity: 28.92% Mismatch.
RESULT 808
ID ADM60974 standard; DNA, 3122 BP.
DE Murine adipocyte specific DNA SeqID 109.
 Indels:
 Indels:
 ADM66974 standard; DNA; 3122 BP.
Murine adipocyte specific DNA SeqID 109.
WO2004011618-A2.
OS-FEB-2004.
 ID AD035561 standard; DNA; 3158 BP.
DE Novel mouse gene sequence #234.
PN W02004046310-A2.
PD 03-UIN-2004.
PA (FIVE-) FIVE PRIME THERAPEUTICS INC.
Percent Similarity: 40.58% Cons Best Local Similarity: 30.43% Mism Query Match:
RESULT 810
 PD 12-DEC-2002.

PA (JOHJ) JOHNSON & JOHNSON RES PTY LTD.

Percent Similarity: 40.45% Cons.

Best Local Similarity: 28.09% Mism
 .v. standard; DNA; 3542 BP. Human bcl-2 gene SEQ ID No 4. WO200299900-Al.
 ACH04172 standard; cDNA; 3717 BP.
 ABL06676 standard; cDNA; 4230 BP.
 PD 21-MAY-2004.
A (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 44.05%
Best Local Similarity: 29.76%
 44.05%
29.76%
13.42%
 40.45%
28.09%
13.42%
 38.27%
28.40%
13.42%
 30.23%
 Percent Similarity: 46.25%
Best Local Similarity: 28.75%
Query Match: 13.42%
RESULT 809
 Percent Similarity: 37.37%
Best Local Similarity: 28.28%
Query Match: 13.42%
RESULT 812
 44.19%
 24-JUN-2004.
(DIAD-) DIADEXUS INC.
 PD 03-APR-2003.
PA (LASE/) LASEK A W.
Percent Similarity: 4
Best Local Similarity: 33
Query Match: 1:
 (HMGE-) HMGENE INC.
 Query Match:
RESULT 814
ID ADP07683 standard;
 Query Match:
RESULT 811
 Query Match:
RESULT 815
 Human secreted protein-encoding gene 7 cDNA clone HKABW11, SEQ ID NO:17. 25-MAY-2001.
 AADO8514 standard; cDNA; 2755 BP.
Human secreted protein-encoding gene 7 cDNA clone HKABW11, SEQ ID NO:37.
WO200136432-A2.
 SEQ ID NO 29425
 Pneumocystis carinii major surface glycoprotein gene HMSG35.
WO200009760-Al.
24-FEB-2000.
 Pneumocystis carinii major surface glycoprotein gene HMSG33 WO200009760-A1.
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16
 cDNA; 2948 BP.
human diagnostic protein #16953.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
 Drosophila melanogaster genomic polynucleotide WO200171042-A2.
 Conservative:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Mismatches:
 Mismatches:
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 ABN95766 standard; DNA; 2336 BP.
Gene #2264 used to diagnose liver cancer.
WO200229103-A2.
 ID AA294069 standard; DNA; 3072 BP.

DE Pneumocystis carinii major surface glycop
N W0200009760-Al.

24-FEB-2000.

PA (USSH) US DEPT HEALTH & HUMAN SERVICES.

Percent Similarity: 50.60% Conserv
 PN WOZ00009760-A1.
PD 24-FEB-2000.
PA (USSH) US DEPT HEALTH & HUMAN SERVICES.
Percent Similarity: 50.60% Consex
Best Local Similarity: 28.92% Mismat
 AAH17443 standard; cDNA; 2825 BP.
Human cDNA sequence SEQ ID NO:16894.
EP1074617-A2.
 AAD08494 standard; cDNA; 2437 BP
 ABL25984 standard; DNA; 2376 BP.
 AAZ94066 standard; DNA; 3084 BP
 ID AAD08494 standard; cDNA; 2437 BDB Human secreted protein-encoding PN WC200136432-A2.
PD 25-MAY-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 40.45%
Best Local Similarity: 28.09%
QUEYY MATCH:
 DNA; 3054 BP
 DE Drosoph.

DE Drosoph.

PN W020011042-Az.,

PD 27-SEP-2001.

PA (PEKE) PE CORP NY.

Percent Similarity: 28.40%

"atch: milarity: 13.42%

"atch: "arch: cDNA; 2'
 EPlure...
07-FEB-2001.
(HELL) HELIX RES INST.
rcent Similarity: 42.86%
ID ARN95766 standard; DNA; 2336
DE Gene #2264 used to diagnose
PN W020029103.A2.
PD 11-ARR-2002.
PA (GENE-) GENE LOGIC INC.
Percent Similarity: 39.33%
Best Local Similarity: 29.21%
Query Match: 13.42%
RESULT 800
 43.53%
28.24%
13.42%
 50.60%
 13.42%
 13.42%
 DE Human CDNA sequence
PN EP1074617-A2.
PD 07-FEB-2001.
PA (HBL1-) HELLY RES IN
Percent Similarity:
Best Local Similarity:
 DNA encoding novel WO200175067-A2.
 AAS81149 standard;
 AAZ94068 standard;
 PA (HYSE-) HYSEQ INC.
Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
Query Match:
RESULT 807
 11-OCT-2001
 Query Match:
 Match:
 Query Match:
RESULT 806
 RESULT 805
```

```
Query Match:
 Query Match:
 RESULT 830
 DE
PN
PD
 promoter contained within a plasmid.
 AAK66749 standard, DNA, 9124 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:21561.
WO200157182-A2.
 SEQ ID NO:21556
 Query Match:

RESULT 820

ID ABL09721 standard; cDNA; 6228 BP.

DE Drosophila melanogaster expressed polynucleotide SEQ ID NO 23645.

PN WO200171042-A2.

PD 27-SEP-2001.
 ABL29886 standard; DNA; 6276 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 41131.
W0200171042-A2.
27-SEP-2001.
(PEKE) PE CORP NY.
 seguence
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 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
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Mismatches:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 AAK66744 standard; DNA; 5079 BP.
Human immune/haematopoietic antigen genomic
WO200157182-A2.
 ABK14253 standard; DNA; 5686 BP.
Human related RAS viral oncogene (RRAS) #2.
WO200188201-A1.
 (RRAS) #1
 Indels:
 Indels:
 Indels:
 ADD67220 standard; DNA; 6183 BP.
Corn Ap1-like (ZM-MADS PRO1) gene
WO2003078590-A2.
 ID ABK14231 Standaru,, DE Human related RAS viral oncogene PW W020018201-A1.
PD 22-NOV-2001.
PA (GENA-) GENAALSSANCE PHARM INC. Percent Similarity: 39.33$

Best Local Similarity: 29.21$

Best Local Similarity: 13.42$
 ABK14231 standard; DNA; 5686 BP.
Human related RAS viral oncogene
WO200188201-A1.
 ID AALLA

DE Human immunc,

PN WO200157182-A2,

PD 09-MCG-2001

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 35-37*

Post Local Similarity: 26.83*

Post Local Similarity: 26.83*
 PA (PION-) PIONEER HI-BRED INT INC. Percent Similarity: 41.05%
Best Local Similarity: 27.37%
 DNA; 9232 BP
 PN W0200188201-A1.
PD 22-NOV-2001.
PA (GENA-) GENAISSANCE PHARM INC.
Percent Similarity: 39.33%
Best Local Similarity: 29.21%
PN WO200171042-A2.
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
Percent Similarity: 38.27%
Best Local Similarity: 28.40%
Query Match: 13.42%
RESULT 816
 Best Local Similarity: 48.48%
Query Match: 13,42%
 Percent Similarity: 44.87%
Best Local Similarity: 26.92%
Ouery Match: 13.42%
 28.41%
13.42%
 13.42%
 40.91%
 ADH69103 standard; D
HIV vaccine V4 DNA.
US2003220276-A1.
27-NOV-2003.
 PA (PEKE) PE CORP NY. Percent Similarity:
 Percent Similarity:
Best Local Similarity:
 (NARA/) NARAYAN O
 25-SEP-2003
 Query Match:
RESULT 822
 Match:
 Query Match
RESULT 819
```

```
Simian-Human immunodeficiency virus clone SHIV-HXBcP 3.2 complete genome. WO200176643-A1. 18-OCT-2001.
 AAD25516 standard; DNA; 10274 BP.
Simian-Human immunodeficiency virus (SHIV) clone 1A11 complete genome.
WO200176643-A1.
 11
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 Conservative:
Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
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 Indels:
 Nucleic acid sequence of SIVmac239 genome.
USSB51813-A.
22-DEC-1998.
 ID AAD25516 standard; D.W., C. Shinan-Human immunodeficiency virus PN WC200176643-A1.
PD 18-OCT-2001.
PPA (BAYU) BAYLOR COLLEGE MEDICINE. Percent Similarity: 28.41% M Best Local Similarity: 28.41% M PATCH.
 AAD25514 standard; DNA; 10006 BP
 AAQ24802 standard; DNA; 10097 BP.
SIVmac239 nef-deletion.
WO9200987-A.
 AAQ22487 standard; DNA; 10279 BP
 AAV81865 standard; DNA; 10279 BP
 (BAYU) BAYLOR COLLEGE MEDICINE.
 BP
 DNA; 9876 BP
 DNA; 9984 BP
PD 09-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 35.37%

Best Local Similarity: 26.83%

Query Match: 13.42%

RESULT 824
 DNA; 9715
 40.91%
28.41%
13.42%
 40.91%
28.41%
 Percent Similarity: 40.91%
Best Local Similarity: 28.41%
Onerv Match: 13.42%
 SIVmac239 proviral genome. WO9200987-A.
 PA (NARA/) NARAYAN O.
Percent Similarity: 40.91%
Best Local Similarity: 28.41%
Query Match: 13.42%
 PD 23-JAN-1992.
PA (HARD) HARVARD COLLEGE.
PECENT Similarity: 40.91%
Best Local Similarity: 28.41%
 PD 23-JAN-1992.
PA (HARD) HARVARD COLLEGE.
Percent Similarity: 40.91%
Best Local Similarity: 28.41%
 13.42%
 13.42%
 13.42%
 PA (NARA/) NARAYAN O.
Percent Similarity: 4
Best Local Similarity: 2
Query Match: 1
RESULT 826
 DE HIV vaccine V7 DNA.
PN US2003220276-A1.
PD 27-NOV-2003.
PA (NARA/) NARAYN O.
Percent Similarity: 2
Query Match:
 ADH69106 standard;
HIV vaccine V6 DNA.
US2003220276-A1.
 HIV vaccine V7 DNA.
US2003220276-A1.
 RESULT 825
ID ADH69104 standard;
 HIV vaccine V5 DNA.
US2003220276-A1.
 ADH69107 standard;
 27-NOV-2003
 27-NOV-2003
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Conservative:

Best Best Query RESUI ID	DE PN PD PACE PECE BEST QUEST TREGUI	DED DED PN PAC PETCE Best Query Query RESUI	ID DE DE PN PA	NE DE PER	PA Perce Best Querr	Perce Best Query	RESUI Perce Best Quest,	Best Query RESUL	010 8 M W W W W W W W W W W W W W W W W W W	Best Query RESUI	P P P P P P P P P P P P P P P P P P P	Best Best Query RESUL ID
11 33 19	11 33 19	11 33 19	8 36 10	ID NO 8573.	6 114 0	ID NO 7801.	6 14 0	ID NO 11245.	6 14 0		16 37 24	541.
Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:	SEQ ID 64. onservative: ismatches: ndels:	9 BP. polynucleotide SEQ I	Conservative: Mismatches: Indels:	SEQ	Conservative: Mismatches: Indels:	BP. polynucleotide SEQ I	Conservative: Mismatches: Indels:		Conservative: Mismatches: Indels:	o. gene, SEQ ID NO:1541
. 2 II - . 3 # % . 4 # %	, DNA, 10372 BP. A. 40.91% 28.41% 13.42%	; DNA; 10535 BP. LLC. 40.918 28.418 13.428	cDNA; 15382 BP a p53 modifier. NC. 43.04% 32.91% 13.42%	DNA; 1681E em related	nowan GENOME SCI INC. larity: 60.00\$ milarity: 42.86\$ 13.42\$	; DNA; 16818 BP. tem related polynucleotide	OME SCI INC. 60.00% 42.86% 13.42%		OME SCI INC. 60.00\$ 42.86\$ 13.42\$	; DNA; 89900 BP. ID 3.	INC. 38.38% 22,22% 13.42%	; DNA; 96597 BP. oma associated ge
Percent Similarity: 40.9 Best Local Similarity: 28.4 Query Match: 13.4 RESULT 83.2	ID ADH69102 standard; DE HIV vaccine V3 DNA PD 27-NOV-2003.  PA (NARA/) NARAYAN O. Percent Similarity: Best Local Similarity: Ouery Match:	ID ADK14025 standard, DN ADK14025 standard, DN SIV genomic DNA. DN US6712612-B1. PN GENE - SO04. PA (GENE - ) GENECURE LLC PA (GENE - ) GENECURE LLC Best Local Similarity: 44 Best Local Similarity: 21 Query Match: 11:	KESULT 834  ID ACD13393 standard; cl DE Human DNA encoding a PN WC200299122-A1. PD 12-DEC-2002. PA (EXEL-) EXELIXIS INC Percent Similarity: 3: Best Local Similarity: 3: Query Match: 11.	KESULT 835  ID ARAIG242 standard; DNA; 16818 BP  NDE Human nervous system related pol  PN WO200159063-A2.  PD IG-AUG-2001.	FA (FUMA-) HUMAN GENCY Percent Similarity: Best Local Similarity: Query Match:	ID ABALS470 standard; DNA; DE Human nervous system rel PN WO200159063-A2.	17 t G	DE Human nervous system related PN W0200159063-A2.	PD 16-AUG-2001. PA (HUMA-) HUMAN GENOME Percent Similarity: 66 Best Local Similarity: 42 Query Match: 02	ID ADO79404 standard; DE DPF3 region, SEQ ID PN WC2004047514-A2.	PA (SEQU-) SEQUENOM INC. Percent Similarity: 38.38* Best Local Similarity: 22.22* Query Match: 13.42*	ALD ADA33023 standard; DNA; 96597 BP DB Mouse Wbnl carcinoma associated PN W02003057146-A2.

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Ty Match:

JLT 847

ABT10748 standard; cDNA; 121600 BP.

Human breast cancer associated coding sequence SEQ ID NO: 882.

WO200259271-A2.
 ЛТ 849
AAP21609 standard; DNA; 349980 BP.
Neisseria meningitidis B nucleotide sequence SEQ ID NO:110.

 #13
 AAA81464 standard; DNA; 102634 BP.
 N. meningitidis partial DNA sequence gnm_12 SEQ ID NO:12.
 20-APR-2000.

 11 848
ApQ18307 standard, DNA; 121600 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 1125.
WO2004048938-A2.
10-UUN-2004.
 9
18
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33
 ADM74618 standard; DNA; 96597 BP.
Murine carcinoma associated (CA) nucleic acid #145.
US2004072154-Al.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
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 Mismatches:
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 Mismatches:
Indels:
 Indels:
 MOUSE Mon1 gene.

WO2003008583-A2.

JO-JAN-2003.

REGAR-) SAGRES DISCOVERY.

RECENT Similarity: 36.00%

REL LOCAL Similarity: 26.67%

MACCH:

ADCESSO3 standard; DNA; 96597 BP.

WOUSE Mon1 genomic sequence.
 (PROT-) PROTEIN DESIGN LABS INC. sent Similarity: 38.20%
: Local Similarity: 30.34%
:y March:
 DNA; 96597 BP
 05-JUN-2003.
(SAGR-) SAGRES DISCOVERY.
rcent Similarity: 36.00%
st Local Similarity: 26.67%
 01-AUG-2002.
(GENE.) GENE LOGIC INC.
rcent Similarity: 38.20%
st Local Similarity: 30.34%
 15-APR-2004.
(MORR/) MORRIS D W.
(ENGE/) ENGELHARD E K.
rcent Similarity: 36.00%
st Local Similarity: 26.67%
ery Match:
36.00%
26.67%
13.42%
 34.94%
24.10%
13.42%
 41.67%
29.17%
13.42%
 34.94%
24.10%
13.42%
 45.56%
23.33%
13.42%
 (CHIR) CHIRON CORP.
cent Similarity: 34
it Local Similarity: 24
iry Match: 13
 LT 840
ADB72761 standard;
 iUIT 844
cent Similarity:
it Local Similarity:
iry Match:
cent Similarity:
 Local Similarity:
cy Match:
 ĹT 845
ent Similarity:
 Local Similarity;
 ent Similarity:
Local Similarity:
 ry Match:
JLT 846
 ry Match:
JLT 842
 IDE ID
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(CURA-) CURAGEN CORP.
 PA (CU
 1D ADD26879 standard; DNA; 429 BP.

DE Human adipocyte Selected Interacting Domain (SID) prey protein DNA #266.

PN WO201290544-A2.

PD 14-NOV-2002.

PA (HYBR-) HYBRIGENICS.

PA (LYNX-) LYNX THERAPEUTICS INC.

Percent Similarity: 44.64 Conservative: 7

Best Local Similarity: 32.14% Mismatches: 28

Query Match: 33.31% Indels: 3
 DE ADDIGESO standard; DNA; 429 BP.

DE ADOIGESO budworm voltage gated calcium channel subunit DNA SeqID 3.

PN WO2004044553-A2.

PD 27-MAY-2004.

PA (FMCC) FMC CORP.

Percent Similarity: 51.06% Mismatches: 22

Query Match: 13.31% Indels: 1
 AAC74694 standard; cDNA; 662 BP.
Human ORFX ORF249 polynucleotide sequence SEQ ID NO:497.
WO200058473-A2.
 33
 12
17
43
 ABN25258 standard; cDNA; 662 BP.
Human ORFX polynucleotide sequence SEQ ID NO:18993.
W0200192523-A2.
06-DEC-2001.
 34
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
Indels:
 ACH79523 standard; DNA; 570 BP.
Human genome derived single exon probe #12718.
US2003194704-A1.
 AAX40600 standard; cDNa; 354 BP.
Human secreted protein 5' EST SEQ ID No: 200.
WO9906550-A2.
 ACH45879 standard; cDNA; 472 BP.
Human foetal brain cDNA #6604.
US2003073623-Al.
 PD 17-APR-2003.

PA (DRAA) DRMANAC R T.

PA (LABA) LABAT I.

PA (STAC/) STACHE-CRAIN B.

PA (DICK/) DICKSON M C.

PA (JOCK/) DICKSON M C.

PA (JOCK/) ASSIGN M C.

PA (JOCK/) ASSIGN M C.

PETCENT Similarity: 40.20%

Best Local Similarity: 33.33%
 PD 16-OCT-2003.
PA (PENN/) PENN S G.
PA (RANK/) RANK D R.
PA (HANZ/) HANZEL D K.
Percent Similarity: 40.59%
Best Local Similarity: 28.71%
Query Match: 13.11%
PN WO200066791-A1.
PD 09-NOV-2000.
PA (CHIR) CHIRON CORP.
PA (GENO-) INST GENOMIC RES.
Percent Similarity: 34.94%
Best Local Similarity: 241.0%
Query Match: 13.42%
 PN mczczcz
PD 05-000.
PA (CURA-) CURAGEN CORP.
Percent Similarity: 45.95%
Best Local Similarity: 32.43%
 48.57%
28.57%
13.31%
 DE Human secreted prote.
PN W0990550-A2.
PD 11-FEB-199.
PA (GBST) GENSET.
Percent Similarity: 4
Best Local Similarity: 2
 Query Match:
RESULT 856
 Query Match:
RESULT 854
 Query Match:
RESULT 851
 RESULT 855
 RESULT 850
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6	
	Mismatches: 29
~	-4
AAH94539 standard; cDNA; 724 B	ъ.
oetal cuna, seu lu no: 5339-A2.	
02-AUG-2001	
(HYSE-) HYSEQ INC. noent Similarity: 37.25	onservative:
Best Local Similarity: 29.41%	hes:
13.31	ndels: 4
3 standard; cDNA; 742 B	ď
Human foetal cDNA, SEQ ID NO:	682.
NC.	
ent Similarity: 37.	Conservative: 8
Best Local Similarity: 29.41%	Mismatches: 21
13.	r
standard; cDNA; 75	
FX ORF614 polynucleot	de sequence SEQ ID NO:1227.
MOZOG DE CO	
AGEN CORP	
rcent Similarity: 43.43	ive: 1
cal Similar	ťΩ
arcn: 13.31 860	n
N24288 standard; cDNA; 756 B	Ф.
nan ORFX polynucleotide s	equence SEQ ID NO:17053.
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Best Local Similarity: 24.24%	he
13.	m
d. DNA: 87	
5,88-ITS2 DNA	sequence.
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NESE	
Fercent Similarity: 41.388 Best Local Similarity: 29 89%	Mismatches: 34
13	
d; DNA; 875 BP	
5.88-ITS2 DNA	sequence.
PN 0838/89//-A.	
ESE	
nt Similarity: 41	ative: 1
Best Local Similarity: 29.89%	ŧD
13	. <del></del> 1
72	
1 rDNA.	
PN US6309840-B1.	
TNECE	
roent Similarity: 41.38%	Conservative: 1
Best Local Similarity: 29.89%	
/ Match: 13	ч
864 AD34331 standard, DNA, 875 B	
Alternative version of Panax	ginseng ITS1-5.8S-ITS2 rDNA.
0-B1.	
(OICE-/ ONLY CHINESE RONG	<u>.</u>

ID AAD23812 standard; DNA; 875 BP. DE Panax quinquefolius ITS1-5.88-ITS2 rDNA. PN US6309840-B1. D 30-OCT-2001. PA (UYCH-) UNIV CHINESE HONG KONG.	Dest Local Similarity: 41.18% Query Match: 13.31% DESTITE 973
SE HONG	5
41.38% 29.89% 13.31%	PD 30-JUL-2002. PA (INCY-) INCYTE GENOMICS INC. Percent Similarity: 58.14* Best Local Similarity: 39.53* Query Match: 13.31*
301	RESULT 874  ID ADBEG65 stand in plants. DE Human cDNA enc. PN BP1308459-A2.
PD 13-MAR-2003. PA (DOWC) DOW CHEM CO. PA (DOWC) DOW AGROSCIENCES LLC. PERCENT Similarity: 46.27\frac{3}{4} Conservative: 10 Best Local Similarity: 31.34\frac{4}{4} Mismatches: 26 Query Match: 13.31\frac{4}{4} Indels: 10	PD 07-MAY-2003 PA (HELLY RES INST. PA (REAA-) RES ASSOC BIOTECHNOLOGY PETCENT SIMILARITY: 42.17\$ BEST LOCAL SIMILARITY: 30.12\$ QUETY MATCH: 13.31\$
SOLT BE7 ABL66283 standard; DNA; 2061 BP. Lung cancer related gene sequence SEQ ID NO:4620. WQ200194629-A2.	RESULT 875  ID AAS/98235 standard; CDNA; 2663 E  DR DNA encoding novel human diagnor  PN W0200175067-A2.
PD 13-DEC-2001. PA (AVAL-) AVALON PHARM. Percent Similarity: 40.59% Conservative: 12 Best Local Similarity: 28.71% Mismatches: 17 Query Match: 13.31% Indels: 43 PRSHT, 968	PD 11-OCT-2001. PA (HYSE-) HYSED INC. Percent Similarity: 47.54% Best Local Similarity: 26.23% Query Match: 13.31%
ID ABL67788 standard; DNA; 2061 BP.  DE Oesophagus cancer related gene sequence SEQ ID NO:6125.  PN W0200194629-A2.  DD 12-DEC 2001	AB22408 standard; cDNA; 2703 DE Human adipose abundant protein PN WO200296355-A2.
PA (AVAL-) AVALON PHARM. Percent Similarity: 40.59% Conservative: 12 Best Local Similarity: 28.71% Mismatches: 17 Query Match: 13.31% Indels: 43 RESULT 869	PA (GETH) GENENTECH INC. Percent Similarity: 45.95% Best Local Similarity: 32.43% Query Match: 13.31%
DE Human cDNA of the invention SEQ ID NO:559.  PN EP1347046-A1.	ID AB224026 standard; cDNA; 2703 DNA BOARD Alpose abundant protein BN WADARDSCAN
St. Ce	PN MOZOGENZOS AZ. PD OS-DEC-2002. PA (GETH ) GENENTECH INC. PECCENT Similarity: 45.95\$ Best Local Similarity: 32.43\$ Query Match: 13.31\$
5 standard, DNA; 2367 BP. oding a Bacillus thuringiensis toxin designated 46-A2.	TENDLI 8/8  ID ABL23365 standard; DNA; 2757 BE DE Drosophila melanogaster genomic PN W0200171042-A2.
PA (MYCO) NYCOGEN CORP.  Percent Similarity: 50.98% Conservative: 5  Best Local Similarity: 41.18% Mismatches: 10  Query Match: 13.31% Indels: 15	PD 27-SEP-2001. PA (PEKE) PE CORP NY. Percent Similarity: 38.18% Best Local Similarity: 23.64% Query Match: 13.31%
AXXB3386 standard; DNA; 2367 BP.  DE Bacillus thuringiensis toxin 185U2(a) encoding DNA.  PN W09933991-A2.	RESULT 879  ID 12550165 standard; DNA; 2774 BF  DE Pichia pastoris Formaldehyde De  PN WO20001829-A2.
0 40 60 50	PD 13-JAN-2000. PA (RESE) PRESEARCH CORP TECHNOLOG Percent Similarity: 54.17% Best Local Similarity: 33.33% Query Match: 13.31%
AAV30289 standard, DNA, 2370 BP. AAV30289 standard, DNA, 2370 BP. Bacillus thuringiensis insecticidal 81F toxin DNA. WC9818912-A2. 07-MAY-1998. (MYCO ) MYCOGEN CORP.	RESOLI'S 80 ID 10024874 standard; DNA; 2999 BP DE HUMAN SOÍT LISSUE SARCOMA-UPTEG PN WOZOOG448938-A2. PD 10-JUN-2004. PA (PROT-) PROTEIN DESIGN LABS INC

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BP.
.n (AAP) 2.7 kb form cDNA sequence.
 BP.
nic polynucleotide SEQ ID NO 21568.
 BP.
n (AAP) 2.7 kb form cDNA sequence.
 BP.
Dehydrogenase 1 (FLD1) gene.
 BP.
egulated DNA - SEQ ID 7694.
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 Conservative: 16
Mismatches: 41
Indels: 27
 BP.
mostic protein #14039.
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 Conservative: :
Mismatches: :
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 (PROT-) PROTEIN DESIGN LABS INC.
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Human disease related procein DNA Sequence Seqibzus.  Housease related procein DNA Sequence Seqibzus.  HOUSEN MAICH:  OLGEN MAICH:  In ARK77432 standard; DNA; 28091 BP.  CORSTO-) OXFORD BIOMEDICA UK LTD.  CORST. ARK77432 standard; DNA; 28091 BP.  DE Human immune/haematopoietic antigen genomic	Hound disease related protein DNA Sequence Sequiplus.  Hound disease related protein DNA Sequence Sequiplus.  Querty Match: 13.31% Mismatches: 26  Querty Match: 13.31% Indels: 22  RESULT 866  (OXFO-) OXFORD BIOMEDICA UK LTD.  DB. Human immune/haematopoietic antigen genomic sequence SEQ
XFO-) OXFORD BIOMEDICA UK LTD. Similarity: 48.94% Conservative: 2 DB Human immune/haematopoietic antigen genomic sequence SEQ	(OXFO-) OXFORD BIOMEDICA UK LTD.  ID AAK77432 standard; DNA; 28091 BP.  Conservative: 2 DB Human immune/haematopoietic antigen genomic sequence SEQ
Similarity: 48.94% Conservative: 2	SImilarity: 48.94% Conservative: 2
Similarity: 48.94* Conservative: 2 DB Human immune/Naematopoietic antigen genomic sequence SEQ	Similarity: 48.94* Conservative: 2 DB Human immune/haematopoietic antigen genomic sequence SEQ
XFO-) UAFORD BIOMEDICA UK LTD. Similarity: 48.94% Conservative: 2	(UXFO-) UAFORD BIOMEDICA UK LTD. rcent Similarity: 48.94% Conservative: 2
XFO-) OXFORD BIOMEDICA UK LTD. Similarity: 48.94% Conservative: 2 DB Human immune/haematopoietic antigen genomic sequence SEQ	(OXFO-) OXFORD BIOMEDICA UK LTD.  ID AAK77432 standard; DNA; 28091 BP.  Conservative: 2 DB Human immune/haematopoietic antigen genomic sequence SEQ
(OXFO-) OXFORD BIOMEDICA UK LTD.  ID AAX77432 standard; DNA; 28091 BP.  DB Human immune/haematopoietic antigen genomic sequence SEQ	(OXFO-) OXFORD BIOMEDICA UK LTD.  TO AAX77432 standard; DNA; 28091 BP.  TCENT Similarity: 48.94% Conservative: 2
ARSOLI 896 ON-POAC. OXOG. ON-POAC. OXOG. ON-POAC. OXOG. OXOG	AESULY 896  ON-PHAR-7-2003.  ON-PHAR 7-32 STANDARD STORE TO TO TO THE PROPERTY OF THE PROPERTY
OG-MAR-2003.  (OKTO).  (OXTO).	06-MAR-2003. (OXFO-) OXFORD BIOMEDICA UK LTD. rcent Similarity: 48.94% Conservative: 2
06-MAR-2003. (OXFO-) OXFORD BIOMEDICA UK LTD. CCENT Similarity: 48.94\$ Conservative: 2 DE Human immune/haematopoietic antigen genomic sequence SEQ	06-MAR-2003. (OXFO-) OXFORD BIOMEDICA UK LTD. rcent Similarity: 48.94% Conservative: 2
06-WAR-2003. (OXPO-) OXFORD BIOMEDICA UK LTD. CORPORT 2 Similarity: 48.94\$ CONSERVATIVE: 2  OSTORE Similarity: 48.94\$  OSTORE Similarity: 48.94\$	06-MAR-2003. (OXFO-) OXFORD BIOMEDICA UK LTD. Conservative: 2  RESULT 896 ID AAK77432 standard; DNA; 28091 BP. DB Human immune/haematopoietic antigen genomic sequence SEQ
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Discrete genomic polymucleotide SEQ ID NO 21565.   DE Human protease gene.	## BESULT 894    Day Abb62371 Standard, DNA; 21784 BP.
## SEGURA 6313 BP.  ## Audo2314 standard; DNA; 21784 BP.  ## DAAD62314 standard; DNA; 21784 BP.  ## Hunan procease gene.  ## Audo23129726-Al.  ## Mismatches: 16  ## Audo23129726-Al.  ## Mismatches: 27  ## Audo23129726-Al.  ## Conservative: 16  ## Audo23129726-Al.  ## Mismatches: 27  ## Audo23129726-Al.  ## Conservative: 16  ## Audo23129726-Al.  ## Conservative: 16  ## Audo23129726-Al.  ## Audo23129726-Al.  ## Conservative: 16  ## Audo23129726-Al.  ## Audo231274-Al.  ##	## SEGURA 6313 BP.  ## And
National Conservative: 16   13.13   13.14   13.14   13.14   13.14   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15   13.15	NY.  19.13 BP.  NY.  NY.  NY.  19.13 BP.  NY.  NY.  NY.  NY.  19.11 894  NOMESCI TO NO 21565.  NY.  19.31 # Indels:  20.32 # Indels:  20.34 # Indels:  20.35 #
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13.314	13.314
27.17\$   Mismatches: 36   Percent Similarity: 63.04\$   Conservative: 10   Query Matches: 13   Percent Similarity: 63.04\$   Conservative: 10   Conservative: 15   Conservative: 16   DE Human Drotease gene.	27.17\$   Mismatches: 36   Percent Similarity: 63.04\$   Conservative: 10   Query Match: 13.11\$   Indels: 17   RESULT 894   Indels: 18
13.34	13.34
Similarity: 4.134   Conservative: 14   Percent Similarity: 2.134   Conservative: 15   Indels: 17   Indels: 17   Indels: 17   Indels: 18   Indels: 19   Indels: 10   Indels:	Similarity: 4.134
Ministry   1.314	Similarity: 2.174   Mismatches: 14   Parameters   Mismatches: 15   Parameters   Mismatches: 16   Parameters   Mismatches: 17   Parameters   Mismatches: 18   Parameters   Mismatches: 19   Parameters   Mismatches: 10   Parameters   Mismatches: 10   Parameters   Mismatches: 10   Parameters   Mismatches: 10   Parameters   Mismatches: 27   Parameters   Mismatches: 27   Parameters   Mismatches: 27   Parameters   Mismatches: 28   Parameters   Mismatches: 28   Parameters   Mismatches: 29   Parameters   Mismatches: 20
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13.314   Indels:   1   Observative   1   Obser	13.314   Indels:   1   REGULT 83   REGULT 83   Each of the conservative:   1   PAD58602 standard, DNA. 17953 BP.
### 13.314 indels: 1   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10.314   10	## SEGURG NO. 5378 BP.  ## SEG
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Percent Similarity: 22.064   Conservative: 16	Second Similarity: 51.6f   Conservative: 16   Percent Similarity: 22.0f   Mismatchee: 18   Percent Similarity: 22.0f   Mismatchee: 19   Percent Similarity: 22.0f   Mismatchee: 19   Percent Similarity: 22.0f   Mismatchee: 22   Percent Similarity: 22.0f   Mismatchee: 24   Percent Similarity: 22.0f   Mismatchee: 25   Percent Similarity: 22.0f   Mismatchee: 26   Percent Similarity: 22.0f   Mismatchee: 27   Percent Similarity: 22.0f   Mismatchee: 27   Percent Similarity: 22.0f   Mismatchee: 27   Percent Similarity: 22.0f   Percent Similarity: 23.0f   Percent Similarity
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### State	### State   DRAY, 5306 BP   PROBLEM Seq1D 5.   PROBLEM SEQ 5.   PROBLE
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### Second Color No. 1935   Conservative: 15   Conservative: 15   Conservative: 15   Conservative: 15   Conservative: 15   Conservative: 15   Conservative: 16   Conservative: 16   Conservative: 16   Conservative: 16   Conservative: 17   Conservative: 16   Conservative: 17   Conservative: 16   Conservative: 17   Conservative: 17   Conservative: 18   Conservative: 18   Conservative: 19   Conservative: 20   Cons	### Standard DNA, 5306 BP  ### Standard DNA, 5308 BP  ### Standard DNA, 5318 BP  ### Standard DNA, 531
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10   20   20   20   20   20   20   20	The following state   10   28   140   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   15
Page 2017-1979   Page	Page 2017-00-12   Page 2017-
Standard   DAM   DAM   Standard   DAM	State   December   D

2 18 2

6 5 5

33 21

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Human respiratory system associated genomic DNA seq id 599.
US2003215893-A1.
 ADA02648 standard; DNA; 32874 BP.
Human TBX21 carcinoma associated gene, SEQ ID NO:1166.
WO2003057146-A2.
 ABK22784 standard; cDNA; 57273 BP.
Human high bone mass (HBM) polynucleotide clone #7.
WC200192891-A2.
W6-DBC-2001.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
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Mismatches:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 мивувоб6 standard; DNA; 66933 BP.
HBM-related clone contig b200e21-h contig4.
WO200292000-A2.
 Indels:
 Indels:
 Indels:
 Indels:
 Indels:
 DE Human HBM gene region b200e21-h_contig4.

DE Human HBM gene region b200e21-h_contig4.

PN W0200177327-A1.

PD 18-0C7-2001.

PA (GENO-) GENOWE THERAPEUTICS CORP.

Percent Similarity: 44.00% Mismat Best Local Similarity: 29.33% Mismat
 PA (GENO-) GENOME THERAPEUTICS CORP.
PA (UVR.) UNIV CREIGHTON SCHOOL MEDICINE.
Percent Similarity: 44.00%
Descriptional Similarity: 29.33%
Misma
 ADESS896 standard; DNA; 32874 BP.
Human TBX21 gene genomic DNA sequence.
WO2003039484-A2.
 EESULT 908

ID AAA28150 standard; DNA; 41684 BP.

DE Human purH gene genomic DNA sequence.

Percent Similarity: 37.93% Conf.

Best Local Similarity: 28.74% Misr
 21-NOV-2002.
(GENO-) GENOME THERAPEUTICS CORP.
(AMHP) WYETH.
 BP.
 ACC45366 standard; DNA; 66933 BP.
Human HBM gene fragment #7.
WO200292764-A2.
 ID ADB72386 standard; DNA; 32874 BP DE Human TBX21 gene.
PN WO2003008583-A2.
PD 30-JAN-2003.
PA (SAGR-) SAGRES DISCOVERY.
Percent Similarity: 53.06%
Best Local Similarity: 13.31%
Query Match:
 INC.
 42.53% 28.74%
 44.00%
29.33%
13.31%
 53.06%
40.82%
13.31%
 PD 15-MAY-2003.

PA (SAGR-) SAGRES DISCOVERY.

Percent Similarity: 53.06%

Best Local Similarity: 40.82%
 29.33%
13.31%
 13.31%
 13.31%
 13,31%
 17-JUL-2003.
(SAGR-) SAGRES DISCOVERY.
 20-NOV-2003.
(HUMA-) HUMAN GENOME SCI
 Local Similarity:
 Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
 DE Human respirator
PN 922003215893-A1.
PD 20-NOV-2003.
PA (HUMA-) HUMAN GE
Percent Similarity:
 PD 21-NOV-2002.
PA (GENO-) GENOME 1
PA (AMHP) WYETH.
Percent Similarity:
 Query Match:
RESULT 911
 Query Match:
 Query Match:
 Query Match:
 Query Match:
 RESULT 906
 Query Matcl
RESULT 907
 RESULT 909
 Human immune/baematopoietic antigen genomic sequence SEQ ID NO:32246.
WO200157182-A2.
 Genomic sequence #5 encoding for novel human respiratory antigen.
 ABA15379 standard; DNA; 28120 BP.

Human nervous system related polynucleotide SEQ 1D NO 7710.

Mo200159663-A2.

16-A00-2001.

(HUMA-) HUMAN GENOME SCI INC.
 1D NO 7706
 AAS30619 standard, DNA, 31994 BP.
DNA encoding novel lung cancer antigen, Seg ID No 71
WO200155300-A2.
 12
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 ABA15375 standard; DNA; 28091 BP.
Human nervous system related polynucleotide SEQ
WC200159063-A2.
 Conservative:
Mismatches:
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Mismatches:
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 Conservative:
Mismatches:
 Conservative:
 Conservative:
 ACA03382 standard; DNA; 31994 BP.
DNA encoding human lung cancer antigen HCLCR09.
US2002173454-A1.
 Mismatches:
 Mismatches:
 Mismatches:
 Indels:
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 Novel lung cancer antigen genomic DNA #1.
US2003049703-A1.
 AAK77434 standard; DNA; 28120 BP
 DNA; 31994 BP
 ADB96730 standard; DNA; 31994 BP.
 ВЪ
 DE Human immune/haematopoietic ant.
PN 9020012182-A2.
PD 09-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 35.46%
Query Match: 13.31%
RESULT 899
 ID AAS30619 standard; DNA; 31994 BD DNA moroding novel lung cancer. PN WO200155300-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 28.74*
Best Local Similarity: 28.74*
 ID A...

DE Human II.C.

PN WO200159063-Az.

PD 16-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI II.C.

Percent Similarity: 35.46%

Rest Local Similarity: 33.96%

Natch: "arch:
 ADG41361 standard; DNA; 31994
 ID ABA15375 standard, DNA, 28091
DE Human nervous system related por wo200159063-A2.
PD 16-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity:
 09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 (HUMA-) HUMAN GENOME SCI INC.
 (HUMA-) HUMAN GENOME SCI INC.
 42.53%
28.74%
13.31%
 36.46%
23.96%
13.31%
 42.53%
28.74%
13.31%
 36.46%
23.96%
13.31%
 42.538
 DE Genomic sequence #5
PN W0200155448-A1.
PD 02-AUG-2001.
PA (HUMA) HUMAN GENOM!
Percent Similarity:
Best Local Similarity:
 ID ADB96730 standard; ID Novel lung cancer ar US2003049703-A1.
PD 13-MAR-2003.
PA (HUMA-) HUMAN GENOMI Percent Similarity:
Best Local Similarity:
 PN WO200157182-A2.
PD 09-AUG-2001.
PA (HUMA-) HUMAN GENOM
Percent Similarity:
Query Match:
RESULT 897
 (BARA/) BARASH S C.
 AAS28165 standard;
 21-NOV-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
 Local Similarity:
 Best Local Similarity:
 ID ACA03382 standar
DE DNA encoding hur
PN US2002173454-A1
PD 21-NOV-2002.
PA (ROSE/) ROSEN C
PA (RUBE/) RUBEN S
PA (BARA/) BARASH S
Percent Similarity:
Best Local Similarity
 Query Match:
RESULT 901
 Query Match:
RESULT 902
 Query Match:
RESULT 903
 Query Match:
RESULT 898
 Query Match:
RESULT 904
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11 38

ABT10718 standard; cDNA; 185371 BP

11	4	11 338 4	4	17	ence.	9 8 8 9 8	1190.	9 228 33	33 8 33 8		9 28 33	ID 1744.	5 24 10
Conservative: Mismatches:	invention #7.	Conservative: Mismatches: Indels:	). Conge <i>r</i> vat i ve		o. genomic DNA sequenc	Conservative: Mismatches: Indels:	gene, SEQ ID NO:	Conservative: Mismatches: Indels:	Conservative: Mismatches: Indels:		Conservative: Mismatches: Indels:	ated DNA - SEQ	Conservative: Mismatches: Indels:
THERAPEUTICS CORP. 44.00\$	5933 BP. 1 to the	THERAPEUTICS CORP. 44.00% -y: 29.33% 13.31%	DNA; 77932 BP. snce for mCG9110 s K.	% % 10.00	96599 BE (PRLR)	2 K. 26.32% 13.31%		COVERY. 35.79% 26.32% 13.31%	DNA, 96599 BP. COVERY. 35.79% 13.31%	rd; DNA; 96599 BP. genomic DNA sequence DISCOVERY		, 1484 oma-up	ign LABS INC. 44.26% 36.07% 13.31%
2002. GENOME WYETH. larity:	5 standard; NA sequence 2015-A2. 2002.	GENOME WYETH. arity: milarit	standard; nomic seque 6558-A1. 003. MORRIS D W. BUGELHARD I	milarity:	standard; olactin re 4383-Al.	MORLIS D W. ENGELHARD E arity: milarity:	if 916 ADAO2672 standard; DNA; 96599 B: Human PRLR carcinoma associated WO2003057146-A2.	003. SAGRES DISC arity: milarity:	standard; LR gene. 8583-A2. 003. SAGRES DISC arity: milarity:	ADE55920 standard; Human PRLR gene gen WO2003039484-A2. 15-MAY-2003. (SAGRE, SAGRES DISC	Percent Similarity: 35.79% Best Local Similarity: 26.32% Query Match: 13.31%	5 Btandard; oft tissue 48938-A2. 2004.	Percent Similarity: Percent Similarity: 44 Best Local Similarity: 36 Query Match: RESULT 920
PD 21-NOV-PA (GENO-)PA (AMHP)Percent Simi	Query March: RESULT 913 ID ADE8243 DE Human D PN WO20029 PD 21-NOV-	PA (C PA (A Percent Best Lo Query A	KESULT 914  ID ADL27149  DE MOUSE GE  PD 20-NOV-2  PA (MORR/)  PA (ENGE/)  PA (ENGE/)	Best La Query I RESULT	ID AAL57703 DE Human pr PN US200306	PA (MORK) PA (ENGE/) PA (ENGE/) Percent Simi Best Local S Query Match:	TO.	PD 1 PA (() Percent Best Lo	TE SOLI 31.  DE Human PR PN W0200300 PD 30-JAN-2 PA (SAGR-) PA (SAGR-) Percent Simil Best Local Si Query Match:	DE DE PN WE HE PO PN WE HE PN WE WE HE PN WE WE HE PN WE	Percen Best L Query	KESOLI 1D A DE H PN W	PA ( Percen Best L Query RESULT

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| The Name Application | The Name Application
Human breast cancer associated coding sequence SEQ ID NO: 852 WO200259271-A2.
 175
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 Query Match:

13.20$

RESULT 927

ID ABZ54861 standard; cDNA; 496 BP.

DE Aspergillus oryzae polynucleotide SEQ ID NO 3974.

PN WO200279476-A1.

PD 10-OCT-2002.
 Conservative:
Mismatches:
Indels:
 (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY. (NARE-) NAT RES INST BREWING. (NORQ.) NAT FOOD RES INST MIN AGRIC. It Similarity: 37.50% Conservative:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 DE Human secreted protein 5' EST, SEQ ID NO: 1284.

DR Human secreted protein 5' EST, SEQ ID NO: 1284.

PN EP1033401-A2.

PD 06-SEP-2000.

PA (GEST) GENSET.

Percent Similarity: 42.55% Conservative.

Best Local Similarity: 26.60% Mismatches:
 ID ABN95539 standard; DNA; 363 BP.
DE Gene #2037 used to diagnose liver cancer.
PN W0200029103-A2.
PD 11-APR-2002.
PA (GENE-) GENE LOGIC INC.
Percent Similarity: 48.00% Mismatches Query Match: 30.00% Mismatches RESULT 926.
 PN WCZUGZON
PD 01-AUG-2002.
PA (GENE) GENE LOGIC INC.
Percent Similarity: 43.43*
Best Local Similarity: 31.31*
 42.55%
26.60%
13.20%
 PA (NAAD-) NAT INST
PA (NARE-) NAT RES
PA (NORQ) NAT FOOD
Percent Similarity:
 Query Match:
RESULT 921
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ery Match: SULT 936 ADI31782 st	Human cDNA #1108. US6607879-B1. 19-AUG-2003. (INCY-) INCYTE CORP. COMBATVATIVE.	13.20%	110 ABZ/1954 standard; DNA; 939 BP.  DE Human G protein coupled receptor coding sequence SEQ ID 165.  PN WO2003000735-A2.  PD 03-JAM-2003.  PD (FRCA).	38.20% 22.47% 13.20%	ID AAQ27091 standard; DNA; 1028 BP.  DE XTY26 probe.  PN W09212262-A1.  PD 23-JUL-1992.  PA (INIW ) INIV WASHINGTON	PA (ADEL-) ADELAIDE MEDICAL CENT WOMEN & CHILDREN. Percent Similarity: 38.03* Conservative: 8 Best Local Similarity: 26.76* Mismatches: 40 Query Match: 13.20* Indels: 4 RESULT 939	ID AAA91542 standard; DNA; 1028 BP. DE 1.0kb PstI fragment from Fragile X syndrome DNA library. PN US642576-B1. PD 05-IIM-2001.	PA (WOME-) WOMEN'S & CHILDREN'S HOSPITAL. PA (UNIW ) UNIV WASHINGTON. Percent Similarity: 38.03% Conservative: 8 Best Local Similarity: 26.76% Mismatches: 40 Query Match: 13.20% Indels: 4	SULT 940 AASO1347 standard; DNA; 1028 BP. Human Fragile X Syndrome 1.0 kb PUS6197500-Bl.	PD 06-MAR-2001. PA (ADEL-) ADELAIDE MEDICAL CENT WOMEN & CHILDREN. Percent Similarity: 38.03* Conservative: 8 Best Local Similarity: 26.76* Mismatches: 40 Query Match: 13.20* Indels: 4	SULT 941 ABZ33699 standard; Human colon tumour WO200283070-A2.	PD 24-OCT-2002. PA (CORI-) CORIXA CORP. Percent Similarity: 42.55\$ Conservative: 15 Best Local Similarity: 26.60\$ Mismatches: 26 Query Match: 13.20\$ Indels: 28	8 standard, cDNA, 1104 BP. tumour associated antigen cDNA.	PD 01-APK-1992. PA (WISTAR INST. Percent Similarity: 42.55\$ Conservative: 15 Best Local Similarity: 26.60\$ Mismatches: 26 Query Match: 13.20\$ Indels: 28	standard; CDNA; 1104 BP. etermined cDNA sequence. 9716-A2. COO1. CORIXA CORP. 121ty: 42.55\$
st Local Similarity: 27.27% Match: 13.20% SULT 928	ID AAK62620 standard; cDNA; 508 BP.  DE Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:7680.  PN W0200157182-A2.  PD 09-AUG-2001.  PA (HUMA-) HUMAN GENOME SCI INC.	rcent Similarity: 44.62% st Local Similarity: 27.69% ery Match: 13.20%	RESULI 927489 standard; cDNA; 731 BP. DE Human ORFX ORF1044 polynucleotide sequence SEQ ID NO:2087. PN W0200058473-A2. PD 05-OCT-2000.	or st	RESULT 930 ID ABN19076 standard, cDNA, 731 BP. DB Human ORFX polynucleotide sequence SEQ ID NO:6629. PN WO200192523-A2. PD 06-DRC-2001.	art ge	ID ABZ33722 standard; cDNA; 732 BP. DB Human colon tumour related cDNA sequence SEQ ID NO:1120. PN WC200283070-A2. PD 24-0CT-2002.	(CORI-) CORIXA CORP. rcent Similarity: 42.55 st Local Similarity: 26.66 ery Match: 13.26 SULT 932	ABD06117 standard; DNA; 765 Pseudomonas aeruginosa polyr US6551795-Bl. 22-APR-2003.	(GENO-) GENOME THERAPBUTICS reent Similarity: 46.58% st Local Similarity: 28.77% sry Match: 13.20% SULT 933	,	(AVAL-) AVALON PHARM. rcent Similarity: 42.55 st Local Similarity: 26.60 ery Match: 13.20 SULT 934	ID ABL65387 standard; DNA; 871 BP.  DE Lung cancer related gene sequence SEQ ID NO:3724.  PN W0200194629-A2.  PD 13-DEC-20011.  PD ANTAL NAVION DEFAULT.	(AVALON FRAKM: crent Similarity: 42.55% bt Local Similarity: 26.60% ary Match: 13.20% SULT 935	ID ABL69501 standard; DNA; 871 BP.  DR Prostate cancer related gene sequence SRQ ID NO:7838.  PN W200194629-A2.  PD 13-DEC-2001.  PA (AVAL-) AVALON PHARM.  Percent Similarity: 42.55% Conservative: 15  Best Local Similarity: 26.60% Mismatches: 34

Best Local Similarity: 26.60% Mismatches: 26  Query Match: 13.20% Indels: 28  RESULT 952  ID ADNO4001 standard; cDNA; 1159 BP.  PN WQ20040208479-A2.  PD 08-ARR-2004.	2 # 2 #	ana DNA fragment SEQ ID NO: 1207	cent Similarity: 42.86% st Local Similarity: 27.47% sry Match: 13.20%		5 # # #	01 0.01 (4	PA (CELL-) CELLZOME AG. Percent Similarity: 52.63% Conservative: 13 Best Local Similarity: 29.82% Mismatches: 22 Query Match: 13.20% Indels: 5 RESULT 956	ID ADK64831 standard; DNA; 1218 BP.  DE Disease treating protein complex-derived gene #1601.  PN EP1338608-A2.	PD 27-AUG-2003. PA (CELL-) CELLZOME AG. Percent Similarity: 52.63% Conservative: 13 Best Local Similarity: 29.82% Mismatches: 22 Query Match: 13.20% Indels: 5	NUT 957 ADI02511 standard; Human cDNA differe; US2003166903-A1.	ary st	standard; teroid-induc 49-Bl.	cce st sry	RESOLA 359 ID Standard; cDNA; 1317 BP. DE Human cancer associated gene sequence SEQ ID NO:193. PN WQ20053350-A1. DD 21-SED-2000	ice Lice
Best Local Similarity:26.60\$Mismatches:26Query Match:13.20\$Indels:28RESULT 94ID ABL6852standard; DNA; 1104 BP.DE Kidney cancer related gene sequence SEQ ID NO:6859.PN WO200194629-A2.PD 13-DEC-2001.	rce st st	ancer related gene se 629-A2.	FA GVALL AVALUA FARM.  PERCENT SIMILARITY: 42.55 Conservative: 15  Best Local Similarity: 26.60\$ Mismatches: 26  Query Match: 13.20\$ Indels: 28	RESULT 946 ID ABL62378 standard; DNA; 1104 BP. DE Colon adenocarcinoma related gene sequence SEQ ID NO:715. PN NO200194629-A2.	PD 13-19C-2011. PD (AVAL-) AVAION PHARM. Percent Similarity: 26-60% Mismatches: 26 Query Match: 13.20% Indels: 28	10	PD 14-FEB-2002. PA (GRNE-) GENE LOGIC INC. PA (NLSB ) JAPAN TOBACCO INC. Percent Similarity: 42.55\$ Conservative: 15 Best Local Similarity: 26.60\$ Mismarches: 26	13.20% Indels: DNA; 1104 BP.	o diagnose liver cancer. C INC. 42.55\$ Conservative:	ty: ard; mour	070-A2. 002. CORIXA CORP. arity: 26.60% Mismatches: milarity: 25.50% Indels:	8 standard; DNA; 1104 BP. nal epithelium/peyer's patc 0852-A2. 2002.	st ery	ID ADN/39767 standard, cDNA, 1159 BP.  DE Cancer/angiogenesis/fibrosis-related nucleic acid, SEQ ID NO:C139.  PN WO2003042661-A2.  PD 22-MAY-2007.	g.

us-09-989-293a-377.rng.spdi

st Local Similarity: 25.81% 27 Match: 13.20% SULT 968	standard; DNA; 2000 BP. e, SEQ ID 5192. 0898-A1. SYNGENTA PARTICIPATIONS AG. arity: 38.57% Conservative: 5	31.43% Mismatches: 13.20% Indels:	; DNA; 2000 BP. Seq ID764 rela	(SYGN ) SYNGENTA PARTICIPATIC CECHT Similarity: 40.00% tt Local Similarity: 26.32% sry Match: 13.20% ULT 970	DE Fungal 2BC gene sequence #73. PN W0200224865-A2. PP 28-MAR-2002. PA (MICR-) MICROBIA INC.	cent Similarity: 45.10% It Local Similarity: 35.29% rry Match: 13.20% SULT 971	ID ACA39934 standard; DNA; 2397 BP.  DE Prokaryotic essential gene #21591.  PN W0200277183-A2.  PD 03-CCT-2002.  PA (ELIT-) ELITRA PHARM INC.	rcent Similarity: 47.37% It Local Similarity: 26.32% ry Match: 13.20% SULT 972	AbJ40188 standard Plant cDNA #1188. US2004016025-A1. 22-JAN-2004.		(KATA/) (KREP/) (PROV/) (RICK/)	Percent Similarity: 50.00% Conservative: 5 Best Local Similarity: 39.13% Mismatches: 21 Query Match: 13.20% Indels: 2	TECULI 97 ADBIO083 standard; DNA; 2688 BP.  DE Alloicoccus otitis antigenic protein encoding DNA SEQ ID NO:4979.  PN WO2003048304-A2.	ity ity	RESULT 97-00-00-00-00-00-00-00-00-00-00-00-00-00	FD 12-UN-2003. PA (AMHP) WYETH HOLDINGS CORP. Percent Similarity: 47.54% Conservative: 10
<pre>it Local Similarity: 26.60% ry Match: 13.20% sULT 960</pre>	ADA14516 standard; cDNA; 1317 BP. Mouse spermatogenesis related cDNA sequence SEQ I WC2003068969-Al. 21-AUG-2003. (NISC-) JAPAN SCI & TECHNOLOGY CORP. snt Similarity: 40.45% Conservative:	Best Local Similarity: 22.47% Mismatches: 37 Query Match: 13.20% Indels: 16 RESULT 961	7 standard; glyceraldeh; 50833-A2. 2004.	PA (UYNC-) UNIV NORTH CAROLINA. PA (EDDY/) EDDY P D E M. Percent Similarity: 40.45 Conservative: 16 Best Local Similarity: 22.47* Mismatches: 37 Query Match: 13.20* Indels: 16	ABOUG186 standard, DNA; 1320 BP. DE Pseudomonas aeruginosa polynucleotide #4790. PN US655195-B1. PD 22-ARF-2003.	PA (GENO-) GENOME THERAPEUTICS CORP. Percent Similarity: 46.58% Conservative: 13 Best Local Similarity: 28.77% Mismatches: 31 Query Match: 13.20% Indels: 8	RESULT 963 ID ABZ33721 standard; cDNA; 1341 BP. DE Human colon tumour related cDNA sequence SEQ ID NO:1119. PN WC200283070-A2. PD 24-OCT-2002.	PA (CORI-) CORIXA CORP.  Percent Similarity: 42.55\$ Conservative: 15 Best Local Similarity: 26 Ouery Match: 13.20\$ Indels: 28	RESULT 964  D AAI69877 standard; cDNA; 1370 BP.  DE Human protein 17 coding sequence.  PN WC20170784-A1.	st st sty	100 standard; DNA; 1557 omonas aeruginosa polyr 1795-B1.	PA (GENO) GENOME THEREPETICS CORP. Percent Similarity: 46.58 Conservative: 13 Best Local Similarity: 28.77% Mismatches: 31 Query Match: 13.20% Indels: 8	AEADS7511 standard; cDNA; 1718 BP.  DE Human enzyme (ENZM) cDNA #23.  PN W02003052075-A2.  PD 26-ITNY-2003.	st st	Tos	PD ZZ-UCI-1998. PA (MILL-) MILIENNIUM BIOTHERAPEUTICS. Percent Similarity: 33.87% Conservative: 10

PA (FARB ) BAYER AG. Percent Similarity: 50.00\$ Conservative: 8 Best Local Similarity: 28.95\$ Mismatches: 19 Query Match: 13.20\$ Indels: 0 RESULT 983 ID ADESE891 standard; DNA; 3376 BP.	Rat gene D14014, SEQ ID W02003016475-A2. 27-FEB-2003. (GEHO ) GEN HOSPITAL COI (FARB ) BAYER AG. (FARB ) BAYER AG. ccent Similarity: 28.99 st Local Similarity: 28.99	9 standard, DNA; 3376 BP. 6 D14014, SEQ ID NO 1731. 16475-A2. GEN HOSPITAL CORP. BAYER AG. 1arity: 13.20\$  Conservative: 111arity: 28.95\$  Mismatches: 111arity: 28.95\$	4 standard, DNA, 3393 BP. reast-specific protein coding se. 53077-A2. 2004. DIADEXUS INC. larity: 37.50% Conse. imilarity: 29.17% Misma	indeis: 4 BP. in coding sequence #24	ar ge	74 BP. ein coding sequence #25	Percent Similarity: 37.50% Conservative: 8 Best Local Similarity: 29.17% Mismatches: 31 Query Match: 13.20% Indels: 29 RESULT 988 Istandard; DNA; 4317 BP. DE P. aeruqinosa virulence qene, VIRIS.	US2004122212-A1. 24-UN-2004. (COSS) COSSON P. (KCHL/) KCHLER T. (BENG/) BENGHEZAL M. (MARC/) MARCHETTI A. (DELD/) DELDEN C V. cent Similarity: 28.77%	13.20% Indels: ; DNA; 4658 BP. ific protein coding sequence #28 INC. 37.50% Conservative: 29.17% Mismatches: 13.20% Indels:
st ery SUL	Str. Str.	SUI SUI	PN W0200218415-A1.  PD 07-MAR-2002.  PA (HUMA-) HUMAN GENOME SCI INC.  Percent Similarity: 43.90% Conservative: 13  Percent Similarity: 28.05% Mismatches: 26  Query Match: 13.20% Indels: 20  RESULT 978  ID ADA40081 standard; CDNA; 3107 BP.	Db mulman sected process consistency construction of the management of the managemen	13.20% Indels: ; cDNA; 3107 BP. otein cDNA #SEQ ID 198.	L L	6 standard; coding human 02994-A2. 2002. HUMAN GENO!	rcent Similarity: 43.908 Ft Local Similarity: 28.038 sry Match: 13.208 SULT 981 ADE55895 standard; DNA; 3 Rat gene D14014, SEQ ID NWO2003016475-A2. 27.FEB-2003. (GEHO ) GEN HOSPITAL CORE	PA         (FARE )         BAYER AG.         Conservative: 8           Percent Similarity: 28.95\$         Mismatches: 19           Query Match: 13.20\$         Indels: 0           RESULT 982         Indels: 0           ID ADES5887 standard; DNA; 3376 BP.         BP.           DE Rat gene D14014, SEQ ID NO 1719.         PN WC2003016475-A2.           PD 27-FEB-2003.         PA (GEHO ) GEN HOSPITAL CORP.

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AAK90888 standard; DNA; 9184 BP.
 27-SEP-2001
 Query Match:
RESULT 999
 Query Match:
RESULT 1001
 Query Match:
RESULT 1003
 Query Match:
RESULT 1000
 Query Match:
RESULT 1006
 DE F. graminearum polyketide synthetase related polynucleotide SEQ ID NO:1.

PN WO2004005522-A2.

PD 15-JAN-2004.

PA (BADI) BASF AG.

Percent Similarity: 41.67$

Conservative: 18

Best Local Similarity: 25.00$

Query Match: 13.20$

The land of
 SEQ ID NO:1417.
 ID NO 1359.
 ABZ74270 standard; DNA; 5158 BP.
Secreted protein gene 215 genomic fragment HNGAM58,
WC200277013-A2.
 8
17
 27 29
 8
31
29
 9 7 7
 Human immune system associated gene SEQ ID NO: 251.
WC200200928-A2.
03-JAN-2002.
 ADP84428 standard; DNA; 5050 BP.
Human breast-specific protein coding seguence #27.
WO2004053077-A2.
 ADA98799 standard; DNA; 5158 BP.
Human secreted protein-related DNA sequence #392
WC2003004623-A2.
 Conservative:
Mismatches:
Indels:
 Human secreted protein encoding genomic DNA SEQ WO200277186-A2.
 Conservative:
Mismatches:
 Conservative:
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 Mismatches:
 Mismatches:
 Mismatches:
Indels:
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 Indels:
 Indel8:
 AAAS4013 standard; DNA; 7131 BP.
6-Methylsalicyclic acid coding sequence.
WO200055340-A1.
 21.SED-2000.
(PION-) PIONBER HI-BRED INT INC.
ent Similarity: 37.65%
Local Similarity: 13.20%
 ABZ67836 standard; DNA; 5158 BP
 ABL32278 standard; DNA; 5198 BP
 PD 03-OCT-2002.

PA (HUMA-) HUMAN GENOME SCI INC.
PErcent Similarity: 54.35%
Best Local Similarity: 36.96%
Query Match: 13.20%
 16-JAN-2003. (HUMA-) HUMAN GENOME SCI INC.
 (HUMA-) HUMAN GENOME SCI INC.
 ID ADP84428 standard; Dun, Colling B. Human breast-specific protein WC2004053077-A2.
PD 24-UW-2004.
PA (DIAD-) DIADEXUS INC.
Percent Similarity: 37.50%
Best Local Similarity: 29.17%
 54.35%
36.96%
13.20%
 54.35%
36.96%
13.20%
 45.45%
27.27%
13.20%
 PN WO200200928-A2,
PD 03-JAN-2002.
PA (EPIG-) EPIGENOMICS AG.
Percent Similarity; 45.459
Best Local Similarity: 27.279
Query Match: 13.208
 DE 6-Methylsalicyclic
PN W0200055340-A1.
PD 21-SEP-2000.
PA (FIGN.) PIONER HI-E
Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
Query Match:
RESULT 993
 Best Local Similarity:
 ID ADA98799 standard
DB Human secreted py
NW Q2003004623-A2.
PD 16-JAN-2003
PA (HUMA-) HUMAN GE)
Percent Similarity:
 03-OCT-2002
 Query Match:
RESULT 998
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:37073.
WO200157182-A2.
 Best Local Similarity: 37.50% Mismatches: 21
Query Match: 13.20% Indels: 1
RESULT 1004 standard; DNA; 31168 BP.
DE DNA encoding novel signal transduction pathway protein, Seq ID 1464.
PN W0200154733-A1.
 Drosophila melanogaster expressed polynucleotide SEQ ID NO 18476.
WO200171042-A2.
 Human digestive system antigen genomic sequence SEQ ID NO: 4464 WO200155314-A2.
 Tr 1003
ABL20384 standard; DNA; 18459 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 12625.
WO200171042-A2.
 ABA07295 standard; DNA; 31168 BP.
Human pancreatic cancer related genomic DNA, SEQ ID NO: 614.
WO200155206-Al.
 3 15 0
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 (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY. (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
 DNA encoding Escherichia coli virulence proteins WO200028038-A2.
 Conservative:
Mismatches:
Indels:
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Mismatches:
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Mismatches:
Indels:
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Mismatches:
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 Indels:
 ABL07998 standard; cDNA; 18445 BP
 BP.
 AAA15186 standard; DNA; 11165 BP
 DE DNA encoding.
PN WO200154733-A1.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 45.65$
Rest Local Similarity: 41.30$
""+Ch:
 ADC87252 standard; DNA; 14562 E
Human GPCR gene SEQ ID NO:1705.
EP1270724-A2.
 T 1000
AAK82261 standard; DNA; 11297
ID AARybusc.

DE Human digestive system.

PN 0220015314-A2.

PD 02-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

Percent Similarity: 45.00%

Percent Similarity: 36.67%
 WOLOUSSON

02-MCD-2001.

(HUMA-) HUMAN GENOME SCI INC.

ent Similarity: 45.65$

: Local Similarity: 41.30$

13.20$
 INC.
 09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 D ABL07998 standard; curry, procession of the Drosophila melanogaster expression of 27-SEP-2001.

PA (PEKE) PE CORP NY.
Percent Similarity: 37.50%
Best Local Similarity: 37.50%
 PN WOZOVOZOBO
PD 18-MAY-2000.
PA (MICR-) MICROSCIENCE LTD.
Percent Similarity: 54.55%
Best Local Similarity: 45.45%
 48.00%
30.67%
13.20%
 54.17%
37.50%
13.20%
 Percent Similarity:
Best Local Similarity:
 (PEKE) PE CORP NY.
 Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
 ID ADC87252 standax
DE Human GPCR gene
PN EP1270724-A2.
PD 02-JAN-2003.
PA (NAAD-) NAT INST
PA (ADSC-) CENT ADV
Percent Similarity:
 Percent Similarity:
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Query Match:
RESULT 1016
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RESULT 1019
 Duery Match:
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 Query Match:
 Query Match:
RESULT 1024
 SEQ ID NO:19572.
 DE Human digestive system antigen genomic sequence SEQ ID NO: 3516.
PN WO200155314-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 45.65$ Conservative: 2
Best Local Similarity: 113.20$ Mismatches: 15
Undels: 10
 Genomic sequence #431 encoding novel human enzyme polypeptide
 AAK64760 standard; DNA; 31168 BP.
Human immune/haematopoietic antigen genomic sequence
WO200157182-A2.
 Human BLR1 carcinoma associated gene, SEQ ID NO:1412.
WO2003057146-A2.
 1180
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 Conservative:
Mismatches:
Indels:
 ADM74489 standard; DNA; 32404 BP.
Human carcinoma associated (CA) nucleic acid #79.
US2004072154-A1.
 Conservative:
Mismatches:
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Mismatches:
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 Indels:
 AAS42115 standard; DNA; 31168 BP.
 ADB94607 standard; DNA; 31168 BP.
Novel human protein DNA #216.
US2002168711-A1.
14-NOV-2002.
 RESULT 1011
ID ADB72632 standard; DNA; 32404 BP
 BP
 T 1007
AAK89940 standard; DNA; 31168 BP.
 ADA02894 standard; DNA; 32404 BP
ADC85373 standard; DNA; 32404
 09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
 PN WO200157182-AZ.
PD 09-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC
Percent Similarity: 45.65%
Best Local Similarity: 41.30%
 Query Match: 13.20%
RESULT 1013
ID ADM74489 standard; DNA, 3240
DE Human carcinoma associated (PN US2004072154.A1.
PD 15-APR-2004.
PA (MORR/) MORRIS D W.
PA (MORR/) MORRIS D W.
PA (ENGE/) ENGELHARD E K.
Percent Similarity: 53.85%
Best Local Similarity: 25.64%
 Mouse Blrl coding sequence. WO2003045230-A2.
 PD 30-CAN-2003.
PA (SAGR-) SAGRES DISCOVERY.
Percent Similarity: 53-85$
Best Local Similarity: 13.20$
 53.85%
25.64%
13.20%
 53.85%
25.64%
13.20%
 Local Similarity: 41.30%
Match: 13.20%
 45.65%
 PD 05-JUN-2003.
PA (SAGR-) SAGRES DISCOVERY.
PA (Indiarity: 53.85%
Best Local Similarity: 25.64%
 (SAGR-) SAGRES DISCOVERY
 (ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
 Human BLR1 gene.
WO2003008583-A2.
 Percent Similarity:
 17-JUL-2003
 Query Match:
RESULT 1009
 Query Match
RESULT 1010
 Query Match
RESULT 1012
 RESULT 1008
 Query M
RESULT
 Best
 DE PER DE
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DE Human immunor/haematopoietic antigen genomic sequence SEQ ID NO:24570.

DE Human immunor/haematopoietic antigen genomic sequence SEQ ID NO:24570.

PN WO200157182-A2.

PD 09-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

PPECCENT Similarity: 38.46% Mismatches: 44

Ouery Match: 13.20% Indels: 4
 Query March:
13.20% Indels: 4
RESULT 1015
ID AAKRét29 standard; DNA; 32986 BP.
DB Human immune/haematopoietic antigen genomic sequence SEQ ID NO:39441.
PN WO200157182-A2.
 DE Human hCG23764 Carcinoma associated gene, SEQ ID NO:1604.

PD 17-JUJ-2003.

PA (SAGR-) SAGRES DISCOVERY.

Percent Similarity: 43.28% Conservative: 8

Best Local Similarity: 31.34% Mismatches: 34

Query Match: 13.20% Indels: 4
 8 4 4
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 Mouse carcinoma-associated (CA) gene TBX21.
US2003099963-A1.
 Indels:
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 Human hCG23764 gene genomic DNA sequence. WO2003053224-A2.
 AAD64732 standard; DNA; 34637 BP.
 ADA66370 standard; DNA; 94531 BP.
 ADB72824 standard; DNA; 94529 BP.
Human hCG23764 gene.
WO2003008583-A2.
 ADA03086 standard; DNA; 94529 BP
 PD 9-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 38.46$
Best Local Similarity: 29.49$
 47.95%
31.51%
13.20%
 PD 29-MAY-2003.
PA (MORK/) MORRISD W.
(MORK) ENGELHARD E K.
PETCENT SIMILATILY: 40.00%
Best Local Similarity: 30.00%
 47.54%
31.15%
13.20%
 43.28%
31.34%
13.20%
 31.51%
 53.70%
33.33%
13.20%
 Percent Similarity: 43.28%
Best Local Similarity: 31.34%
Query Match: 13.20%
 13.20%
 47.95%
 PD 03-JUL-2003.
PA (SAGR-) SAGRES DISCOVERY.
Percent Similarity: 43.289
Best Local Similarity: 31.349
 30-JAN-2003.
(SAGR-) SAGRES DISCOVERY
 RESULT 1021
Percent Similarity:
Best Local Similarity:
 RESULT 1022
Percent Similarity:
Best Local Similarity:
 RESULT 1023
Percent Similarity:
Best Local Similarity:
 Best Local Similarity:
 RESULT 1020
Percent Similarity:
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st Local S SULT LO3C SULT 1032 ACH2061 Human a US-2C (DRMA/) (STABA/) (STABA/)	(JONE/) ccent Simi st Local S six sry Match: srlr 1033 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 ADJ4444 (BUDW/) (RNIG/) (GIAZ/)	PA (GOFF) GOFF S.A.  PA (KREPA) KATAGTRI F.  PA (KREPA) KREPS J.  PA (KREPA) RATAGTRI F.  PA (RICK) RICKE D.  PA (ZHUT/) ZHU T.  PETCENT Similarity: 53.57% Conservative: 9  Best Local Similarity: 37.50% Mismatches: 22  Query Match: 13.10% Indels: 4  RESULT 1034  ID ACH20641 standard; CDNA; 411 BP.  DE Human adult liver CDNA #253.  PN US2003073623.A1.  PD 17.APR-2003.  PA (DRAAA) DRMANAR R T.	PA (STAC/) STACHE-CRAIN B.  PA (DICKSON M C.  PA (DICKSON M C.  PA (JONES) JONES L W.  Percent Similarity: 44.44 Mismatches: 15  Best Local Similarity: 27.78 Mismatches: 34  Query Match: 13.104 Indels: 16  RESULT 1035  ID AASGOGOS tandard; CDNA, 431 BP.  DE Human cancer agent-resistance marker #360.  PN WO200179556-A2.  PD 25-CCT-2001.  PA (MILLENNIUM PREDICTIVE MEDICINE INC.  Percent Similarity: 27.784 Mismatches: 15  Best Local Similarity: 27.784 Mismatches: 16  Ouery Match: 13.104 Indels: 16	Conservative: Mismatches: Indels: EST, SEQ ID NO: 154.
DE Ovary cancer related gene sequence SEQ ID NO:6459.  PN W0200194629-A2.  PD 13-DEC-2001.  PP (AVAL.) AVALON PHARM.  Percent Similarity: 29.55*  RESULT 1025  DD HUMAN SOft tissue sarcoma-upregulated DNA - SEQ ID 2392.	PD 10-JUN-2004. PD 10-JUN-2004. PD 10-JUN-2004. PA (PROT-) PROTEIN DESIGN LABS INC. CONSERVATIVE: 31  Best Local Similarity: 29.55\$ Mismatches: 31  Query Match: 13.20\$ Indels: 20  RESULT 10.26  DE MSI-H Carcinoma genomic DNA, 262090 BP. DE MSI-H Carcinoma genomic DNA sequence SEQ ID NO:44.  PA (XIMH/) KIM H G.  PA (KIMH/) KIM H G.  PA (KIMH/) KIM N G.  PA (KIRAL) IRE J S.	rce st sur sur sur	DE Human ORFX ORF300 polynucleotide sequence SEQ ID NO:599.  PN W0200058473-A2.  PD 05-OCT-2000.  PA (UTRA-) CURAGEN CORP.  Percent Similarity: 39.73* Conservative: 7  Best Local Similarity: 30.14* Mismatches: 7  Query Match: 13.10* Indels: 7  RESULT 1029  Human ORFX polynucleotide sequence SEQ ID NO:17435.  PD Human ORFX polynucleotide sequence SEQ ID NO:17435.  PD 06-DEC-2001.  PA (CURA-) CURAGEN CORP.  PA (CURA-) CURAGEN CORP.  Research Similarity: 39.73* Conservative: 7  Best Local Similarity: 30.14* Mismatches: 37	Match: T 1030  AA19280 standard; CDNA; 347 BP. Human polynuclectide SEQ ID NO 10340.  WO200164835-A2. 07-SEP-2001. (HYSE-) HYSEQ INC. Tocal Similarity: 38.89\$ Mismatches: T 1031 AACO157 standard; CDNA; 363 BP. Human secreted protein 5' EST, SEQ ID NO: 155. EP103401-A2. 06-SEP-2000. (GSEP-2000.

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ADE82099 standard; cDNA; 483 BP.
Arabidopsis thaliana expressed polynucleotide seg id 870.
US2003115639-A1.
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IN UGS2003115639-A1.

PA (GORL/) GORLACH J.

PA (GORL/) GORLACH J.

PA (HANY/) AN Y.

PA (HANY/) HAMILTON C M.

PA (RAINY) RAINES T M.

PA (RAINY) RAINES M.

PA (MATH/) MATHEW A V.

PA (RAINY) RAINES W.

PA (MATH/) MATHEW A V.

PA (RAINY) RAINES W.

PA (RAINY) RAINES W.

PA (RAINY) RAINES W.

PA (HOFF) HOFPMAN P.

PA (HOFF) PANIS W.

PA (HANZ) PANIS G.

PA (HANZ) PANIS G.

PA (HANZ) PANIS C.

 Conservative:
Mismatches:
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Mismatches:
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 ACH76127 standard; DNA; 506 BP.
Human genome derived single exon probe #9322.
US2003194704-A1.
 Best Local Similarity: 33.33* Mismatc.
Query Match: 13.10* Indels:
RESULT 1042
DE ABX56971 standard; DNA; 567 BP.
DB Arabidopsis thaliana polynucleotide #323.
PN US2002040409-A1.
PD 04-ARR-2002.
PA (GORL/) GORLACH J.
 T 1038
ACH28025 standard; cDNA; 475 BP
 Query Match:
RESULT 1041
ID ACH7612 standard; DNA; 506 BP DE Human genome derived single ex PN US2003194704-A1.
PD 16-007-2003.
PA (PENN/) PENN S G.
PA (RANK/) RANK D R.
PA (HANZ) HANZEL D X.
PA (HANZEL D X.
PECCENT SIMILATITY: 33.33%
Best Local Similarity: 13.10%
 PN 16-OCT-2003.
PA (PENN/) PENN S G.
PA (HANK/) RANK D R.
PA (HANX/) HANZEL D K.
Percent Similarity: 50 79%
Best Local Similarity: 31.75%
 Human adult ovary cDNA #6405.
US2003073623-A1.
17-APR-2003.
PD 06-SEP-2000.

PA (GEST) GENNET.
Percent Similarity: 44.44%
Best Local Similarity: 27.78%
Query Match: 13.10%
RESULT 103%
ID ACH28025 standard; CDNA; 475
DE Human adult covary CDNA #6405
PW 1052030376523-A1.
PD 17-APR-2003.
PA (LABA/) LABAT I.
PA (LABA/) LABAT I.
PA (STAC/) STACHE-CRAIN B.
PA (JONE/) JONES L W.
PA (JONE/) JONES L W.
Percent Similarity: 25.27%
Query Match: 13.10%
 Query Match:
RESULT 1039
```

04-APR-2002. (GORL/) GORLACH J.

```
ADK53744 standard; DNA; 574 BP.
Plant DNA sequence which confers altered metabolic characteristic #1127.
WO2003020936-A1.
 ADK55700 standard; DNA; 657 BP.
Plant DNA sequence which confers altered metabolic characteristic #3083.
WO2003020936-A1.
 ADK57820 standard; DNA; 657 BP. Plant DNA sequence which confers altered metabolic characteristic #5203. WO2003020936-Al.
 Query Match:
RESULT 1047
RESULT 1047
DB AAK79094 standard; DNA; 659 BP.
DB Human immine/haematopoietic antigen genomic sequence SEQ ID NO:33906.
PN WO200157182-A2.
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 Conservative:
Mismatches:
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Mismatches:
Indels:
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 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 ACH67680 standard; DNA; 595 BP.
Human genome derived single exon probe #875.
US2003194704-Al.
 Indels:
 Indels:
 RESULT 1048
ID ACF66493 standard; DNA; 693 BP
 PD 13-MAR-2003.

PA (DOWC) DOW CHEM CO.

PA (DOW C) DOW AGROSCIENCES LLC.

Percent Similarity: 44.94%

Best Local Similarity: 26.97%

Query Match: 13.10%
 PN MARK-2003.
PD A. (DOWC) DOW CHEM CO.
PA (DOWC) DOW AGROSCIENCES LLC.
Percent Similarity: 44.94*
Best Local Similarity: 26.97*
 (DOWC) DOW CHEM CO. (DOWC) DOW AGROSCIENCES LLC.
 PA (HUMA-) HUMAN GENOME SCI INC.
Percent Similarity: 43.66%
Est Local Similarity: 32.39%
Query Match: 13.10%
PA (ANYY/) AN Y.
PA (HAMI/) HAMILTON C M.
PA (PRIC/) PRICE J L.
PA (RAIN/) TAINES T M.
PA (YUYY/) YU Y.
PA (RAME/) RAMEAKA J G.
PA (RAME/) RAMEAKA J G.
PA (MATH/) MATHEW A V.
PA (LEDF/) LEDFORD B L.
PA (MOES/) WOESENER J P.
PA (HAS/) HAAS W D.
PA (GARC/) GARCIA C A.
PA (GARC/) GARCIA C A.
PA (RRIC/) RITCKER M.
PA (RIC/) RITCKER M.
PA (RIC/) ALLEN Y.
PA (HURB/) HURBAN N.
PA (HURB/) HOFFMAN N.
PA (HURB/) HOFFMAN N.
PA (HURB/) HURBAN P.
PECCENT SIMILATIFY:
PA (HURB/) HOFFMAN N.
PECCENT SIMILATIFY:
PECCEN
 ID ADK53744 standard; DNA; 574
DE Plant DNA sequence which cor
PN W0200320936-A1.
PD 13-MAR-2003.
PA (DOWC) DOW CHEM CO.
PA (DOWC) DOW AGROSCIENCES LLC
Percent Similarity: 33.338
Query Match: 13.108
 ent Similarity: 40.30%
Local Similarity: 31.34%
Match: 13.10%
 16-OCT-2003.
(PENN/) PENN S G.
(RANK/) RANK D R.
(HANZ/) HANZEL D K.
 ID ACH67680 standar
DE Human genome der
PD 16-OCT-2003.
PA (RANK) PENN S G
RANK) RANK D R
PA (HANZ) HANZEL D
PERCENT Similarity:
 09-AUG-2001
 Query Match:
 RESULT 1045
 RESULT 1046
 RESULT 1043
```

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24
 13
34
11
 9 2 2
 12
25
7
 3 20
 942
 1141
 Percent Similarity: 47.27$ Conservative: 9 Best Local Similarity: 30.91$ Mismatches: 24 Query Match: 13.10$ Indels: 5 RESULT 1057
ID AAAC45528 standard; DNA; 1024 BP.
DB Arabidopsis thaliana DNA fragment SEQ ID NO: 46825.
 AAC41895 standard; DNA; 1024 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 33532.
EP1033405-A2.
 Percent Similarity: 47.27$ Conservative: 9
Best Local Similarity: 30.91$ Mismatches: 24
Query Match: 13.10$ Indels: 5
RESULT 1058
ID AASB3801 standard; cDNA; 1036 BP.
DE DNA encoding novel human diagnostic protein #19605.
 Conservative:
Mismatches:
Indels:
 Conservative:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
 Mismatches:
Indels:
 Mismatches:
Indels:
 Mismatches:
 Indels:
 Indels:
 PD 11-CCT-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity: 46.67% Conservational Similarity: 26.67% Mismat Query Match: 13.10% Indelto ID AAF07696 standard; CDNA; 1124 BP.
DE RUSALIM Venematum EST SEQ ID NO:219.
 BP.
ID 2577.
 PD 24-070-2003.
PD 64-070-2003.
PA (GENO-) GENOME THERAPEUTICS CORP.
Percent Similarity: 45.78*
Best Local Similarity: 30.12*
Acch. 13.10*
 Percent Similarity: 41.03 $
Best Local Similarity: 26.92 $
Query Match: 13.10 $
RESULT 1060
ID ADJ39371 standard; cDNA, 1158 BP.
 28-SEP-2000.
(NOVO) NOVO NORDISK BIOTECH INC.
(NOVO) NOVO NORDISK AS.
 ADC92950 standard; DNA; 996
E. faecium DNA sequence SEQ
US6583275-B1.
 47.27%
30.91%
13.10%
 48.89%
40.00%
13.10%
 BRIGGS S P.
COOPER B.
GLAZEBROOK J.
GOFF S A.
KATAGIRI F.
 BUDWORTH P. MOUGHAMER T.
 PA (ZHUT/) ZHU T.
Percent Similarity;
Query Match:
RESULT 1061
 SLATER T.
DAVIS K R.
ALLEN K.
HOFFMAN N.
 KREPS J.
PROVART N.
 Best Local Similarity:
 KRICKER M.
 (HURB/) HURBAN P.
 Plant cDNA #371.
US2004016025-A1.
 PA (KRIC/) KRICKER
PA (SLAT) SLATER TE
PA (DAVI/) DAVIS K
PA (ALLE/) ALLEN K.
PA (HORF/) HOFFMAN
PA (HURBA) PERCENT SIMILALY:
 RICKE D.
 06-SEP-2000.
 06-SEP-2000
 22-JAN-2004
(BUDW/) BUD
 (GLAZ/)
(GOFF/)
(KATA/)
 Query Match:
RESULT 1056
 (BRIG/)
(COOP/)
 Query Match:
RESULT 1055
 (MOUG/)
 (KREP/)
 (PROV/)
 (RICK/)
 DE FILC.

PN WO20029486,

PN WO20029486,

PA (CNRS) INST PASTEUR.

PA (CNRS) CNRS CENT NAT RECH SCI.

PA (CNRS) CNRS CENT NAT RECH SCI.

Percent Similarity: 26.92$ Mismatches: 32

CHEST AAH92909 standard; DNA, 700 BP.

DD AAAH92909 standard; DNA, 700 BP.

DE Human inflammatory bowel disease related gene fragment IGR1375a.

"TN-2001."

"TN-2001."

"TN-2001."

"TN-2001."

"TN-2001."

"TN-PRECUTICS CORP.

CONSERVATIVE: 6

CONSERVATIVE: 6
 Staphylococcus aureus DNA for cellular proliferation protein #1470. W0200170955-A2. 27-SEP-2001.
 AAS52223 standard; DNA; 789 BP.
Staphylococcus aureus DNA for cellular proliferation protein #640.
WO200170955-A2.
 ABN98574 standard; DNA; 949 BP.
Arabidopsis thaliana expressed polynucleotide SEQ ID NO 342.
US2002023281-A1.
 20
26
43
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 (INWR) MERIEUX ORAVAX PASTEUR MERIEUX SERUMS.
(HUMA-) HUMAN GENOME SCI INC.
 AAH07926 standard; cDNA; 825 BP.
Human cDNA clone (5'-primer) SEQ ID NO:4761.
EP1074617-A2.
 Indels:
 Indels:
 Indels:
 Indel8:
 RESULT 1050
TD AAX13949 standard; DNA; 766 BP.
DE H. PYLOTI GHPO 1578 gene.
PN WO9843478-A1.
 RESULT 1052
ID AAS55158 standard, DNA, 789 BP
 PN WO9843478-A1.
PD 08-OCT-1998.
PA (INMR) MERLEUX ORAVAX PASTEP PA (HUMA-) HUMAN GENOME SCI INV Percent Similarity: 48.05% Best Local Similarity: 27.27%
 PD 27-SED-2001.
PA (SLIT-) ELITRA PHARM INC.
Percent Similarity: 41.03%
Best Local Similarity: 13.93%
 PN DETECTION OF THE PROPERTY OF THE PROPERTY OF THE PROPERTY AND THE PROPERTY OF THE PROPERTY
 Best Local Similarity: 41.03% Query Match: 13.10%
 (ELIT-) ELITRA PHARM INC
 AN Y.
HAMILION C M.
PRICE J L.
RAINES T M.
YU Y.
 WOESSNER J P.
HAAS W D.
GARCIA C A.
 RAMEAKA J G.
 MATHEW A V.
LEDFORD B L.
 21-FEB-2002.
(GORL/) GORLACH J.
 PAGE A.
 (HAMI/)
(PRIC/)
(RAIN/)
 (WOES/)
(HAAS/)
(GARC/)
 Query Match:
RESULT 1051
 (RAME/)
(PAGE/)
 ANYY/)
 Query Match:
RESULT 1054
 XUXX/)
 MATH/)
 LEDF/)
```

```
04-SEP-2003
 28-MAR-2002
 09-AUG-2001
 Query Match:
RESULT 1070
 Query Match:
 Query Match:
 Query Match:
 Query Match:
 RESULT 1072
 RESULT
 RESULT
 ABQ45449 standard; DNA; 1283 BP. Oligonucleotide for detecting cytosine methylation SEQ ID NO 32040.
WO200218632-A2.
 Oligonucleotide for detecting cytosine methylation SEQ ID NO 32039. WO200218632-A2.
 Drosophila melanogaster expressed polynucleotide SEQ ID NO 33413. WO200171042-A2.
 ADE76867 standard; cDNA; 1629 BP.
Human cDNA differentially expressed in a liver disorder #22.
US2003108871-A1.
 Query Match:
RESULT 1065
ID ADC1334 standard; cDNA; 1411 BP.
DE Human NOVX protein encoding cDNA sequence, SEQ ID No 13.
PN W02003004617-A2.
 10
 9
5
 26
 7
22
55
 32
 8
13
39
 cDNA; 1188 BP.
human diagnostic protein #16847.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indel8:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Mismatches:
Indels:
 Indels:
 Indels:
 Prokaryotic essential gene #29503.
WO200277183-A2.
 Query Matcn:
RESULT 1066
ID ABL12977 standard; cDNA; 1521 BP.
 ABQ45448 standard; DNA; 1283 BP.
 ADES8172 standard; DNA; 1674 BP.
Rat gene M91466, SEQ ID NO 4043.
W02003016475-A2.
27-FEB-2003.
 ACA47846 standard; DNA; 1394 BP
 ID ABQ45448 standaru,,
DE Oligonucleotide for detecting
DN W0200218632-A2.
PD W7-MAR-2002.
PA (EPIG-) EPIGENOMICS AG.
Percent Similarity: 61.54*
Best Local Similarity: 38.46*
 (GEHO) GEN HOSPITAL CORP. (FARB) BAYER AG.
 43.64%
34.55%
13.10%
 PA (PEKE) PE COLP NY.

Percent Similarity: 41.46%
Best Local Similarity: 23.17%
Query Match: 13.10%
RESULT 1067
 44.44%
27.78%
13.10%
 33.04%
26.96%
13.10%
 Percent Similarity: 61.54%
Best Local Similarity: 38.46%
Query Match: 13.10%
 PA (ELIT-) ELITRA PHARM INC.
Percent Similarity: 49.28%
Best Local Similarity: 18.84%
Ouery Match: 13.10%
 Percent Similarity: 37.35%
Best Local Similarity: 27.71%
Query Match: 13.10%
 07-MAR-2002.
(EPIG-) EPIGENOMICS AG
ID AASB1043 standard; cl
D DNA encoding novel hr
PN WC200175067-A2.
PD 11-OCT-2001.
PA (HYSE) HYSEQ INC.
Percent Similarity: 3:
 Local Similarity: Match:
 Best Local Similarity:
Query Match:
RESULT 1069
 PA (KASE/) KASER M R. Percent Similarity:
 PA (FARB) BAYER /
Percent Similarity:
 27-SEP-2001.
 03-OCT-2002
 12-JUN-2003
 Query Match:
RESULT 1063
 Match:
 Query Match
RESULT 1068
 Query Match
RESULT 1064
 Query M
RESULT
 Best
 DE PER
 PN
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ID ACF39414 standard; DNA; 1737 BP.

DB Mycobacterium tuberculosis mycobacterial antigen DNA SEQ ID NO:206.

PN WOOD333530-A2.

PD 24-APR-2003.

PA (MICR-) MICROBIOLOGICAL RES AUTHORITY.

PRECENT Similarity: 47.92% Conservative: 4

Best Local Similarity: 39.58% Mismatches: 23

Query Match: 13.10% Indels: 2
 Human fibrinogen, B beta polypeptide (FGB) DNA sequence. WO2003072827-A1.
 Aurese31 standard; DNA; 1918 BP.
Human fibrinogen, B beta polypeptide (FGB) DNA sequence.
W0200172827-A1.
 34
34
16
 34
15
 34
16
 15
34
16
 Human fibrinogen beta-chain mRNA, partial cds DNA.
 Human fibrinogen beta-chain mRNA, partial cds DNA.
WO2003072827-A1.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
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Mismatches:
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Mismatches:
 Conservative:
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 Mismatches:
 Mismatches:
 Indels:
 Indels:
 Indels:
 PN WO2003072827-A1.
PD 04-SEP-2003.
PA CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.
PA CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.
BACCENT SIMILATILY: 44.44*
Mismatc
 PA (CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.
Percent Similarity: 44.44* Conserv
Best Local Similarity: 27.78* Mismatc
 PA (CHIL.) CHILDREN'S HOSPITAL MEDICAL CENT.
Percent Similarity: 44.44% Conserv
Best Local Similarity: 27.79% Mismatc
 (CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT
 Human polynucleotide SEQ ID NO 170. 90-AMT.
 Human polynucleotide SEQ ID NO 171. WO200157190-A2.
 RESULT 1077
ID AADS7334 standard; cDNA; 2024 BP
 RESULT 1073
ID ADP65102 standard; DNA; 1918 BP.
 ABN79808 standard; DNA; 1962 BP.
 ADP65552 standard; DNA; 1883 BP
 AAK51626 standard; cDNA; 1862
 T 1071
ADP65676 standard; DNA; 1883
 Fungal ZBC gene sequence #45.
WO200224865-A2.
 PD 09-AUG-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity: 27.78%
Best Local Similarity: 27.78%
 44.44%
27.78%
13.10%
 44.44%
27.78%
13.10%
 44.44%
27.78%
13.10%
 44.44%
27.78%
13.10%
 Percent Similarity: 44.44%
Best Local Similarity: 27.78%
Query Match: 13.10%
 47.318
 PA (MICR-) MICROBIA INC.
Percent Similarity: 47
Best Local Similarity: 32
Query Match: 13
 RESULT 1074
ID ADP65031 standard;
 PA (HYSE-) HYSEQ INC.
Percent Similarity:
Best Local Similarity:
 DE Human fibrinogen
PN W02003072837-A1.
PD 04-SEP-2003.
PA (CHIL-) CHILDREN
Percent Similarity:
 04-SEP-2003
```

Percent Similarity: 12.513 Best Local Similarity: 38.468 Query Match: 13.108	. 025	c	HUMAN GE	
	Conservative: Mismatches: Indels:	11 19		Conservativ Mismatches
ADAS3914 standard; cDNA; 2033 BP Human coding sequence, SEQ ID 14	82.		Query match: RESULT 1086 ID ADA98091 standard; cDNA; 2186 BP	
PN BF1293569-AZ. PD 19-MAR-2003. PA (HELL-) HELIX RES INST.			D C (1	ence #185.
PA (REAS-) RES ASSOC BIOTECHNOLOGY. Percent Similarity: 39.08%	Conservative:	6.6	Ō	Conservati
Best Local Similarity: 28.74% Query Match: 13.10% RESH.T 1079	Mismatches: Indels:	22 31	Best Local Similarity: 35.90% Query Match: 13.10%	Mısmatches Indels:
4 standard; DNA; 2070 BP. psis thaliana DNA fragmen 05-82	t SEQ ID NO: 37710	10.	ID ADA4396 standard; cDNA; 2186 BP. DE Human secreted protein cDNA SEQ ID PN WOOD3000865-A2	ID 154.
PD 06-SEP-2000. Percent Similarity: 49.06*	Conservative:	9	PD 03-JAN-2003. PD 03-JAN-2003. PA (HUMA-) HUMAN GENOME SCI INC.	
Desc Docal Similarity: 32.00%		10	Best Local Similarity: 35.90%	Mismatches
KESULT 1080 ID ABZ13390 standard; DNA; 2070 BP. DE Arabidopsis thaliana stress regulated DN MOTORO16665.20	gene SEQ	ID NO 1195,	86 E	Indels: IP.
002. SCRIPPS RES INST.	Ç		787-A2. 002. TIMAN CENOME SCT INC	)    -
arity: 49.06% milarity: 32.08%	Conservative: Mismatches: Indels:	9 17 10 10 10 10 10 10 10 10 10 10 10 10 10	arity: 46.15% nilarity: 35.90%	Conservativ Mismatches
RESULT 1081 TD AD851220 Standard: DNA: 2080 RP		24	at and and.	
Primary rat hepatocyte toxicity W02003065993-A2.	modelling relate	ed gene SEQ ID NO:3762.	ing.	sequence #9
Ŋ	. 011:11	a	002. HUMAN GENON	3
Concern Similarity: 30.43% Query Match: 13.10%	Mismatches: Indels:	11 29	Fercent Similarity: %0.15% Pest Local Similarity: 35.90% Ouery Match: 13.10%	Mismatches Indels:
ULT 1082 AAT10252 standard; cDNA; 2085 BP		•	ij	
DE Human calpastatin coding sequence PN W09533060-A1. PD 07-DEC-1995.	•		DE Arabidopsis thaliana DNA fragment PN EP1033405-A2. PD 06-5FP-2000.	t SEQ ID NO
E E	Conservative:	12	Percent Similarity: 49.068 Best Local Similarity: 32.088	Conservativ
Best Local Similarity: 26.67% Query Match: 13.10%	Mismatches: Indels:	25 7	13.10%	Indels:
1; DNA; 2148 BP.			ID ADL61097 standard; DNA; 2290 BP. DE Human protein tyrosine kinase bi	BP. • biomarker calpa
carbascacin-i	ike polypeptide.		M02004020583-AZ.	
(TAKI ) TAKARA SHUS	4	,	S i	Conservativ
Query Match: 13.10%	Mismatches: Indels:	12 25 7	Query Match: 13.10% RESULT 1092	Indels:
8 standard; nzyme ENZM-9			ID AAS03910 standard, cDNA, 2450 BP. DE Human secreted protein gene #29. PN WO200123598-A1.	
WO2003042357-A2. 22-MAY-2003.				
PA (INCY-) INCYTE GENOMICS INC. Percent Similarity: 46.15% Best Local Similarity: 35.90%	Conservative: Mismatches:	88 9.4 9.5	Percent Similarity: 40.91% Best Local Similarity: 25.00% Query Match: 13.10%	Conservativ Mismatches: Indels:
9 standard;		Q 4		80 cDNA clone

```
ABZ73460 standard; cDNA; 2450 BP.
Secreted protein-encoding gene 180 cDNA clone HLQAS12, SEQ ID NO:190.
 DNA; 2290 BP.
sine kinase biomarker calpastatin DNA.
 8
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17
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23
 12
25
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 14
39
13
 DNA; 2244 BP.
.na DNA fragment SEQ ID NO: 38085.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
Indels:
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Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 DNA; 2186 BP.
 : DNA; 2186 BP.
stein coding sequence #188.
 tein cDNA sequence #185.
 cDNA; 2186 BP.
tein cDNA SEQ ID 154.
 CDNA; 2186 BP.
 YERS SQUIBB CO.
46.67%
26.67%
13.10%
 cDNA; 2450 BP.
 OME SCI INC.
46.15%
35.90%
13.10%
 OME SCI INC.
46.15%
35.90%
13.10%
 OME SCI INC.
46.15%
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13.10%
 OME SCI INC.
46.15%
35.90%
13.10%
 OME SCI INC.
40.918
25.008
13.108
ME SCI INC
 46.15%
35.90%
13.10%
 49.06%
32.08%
13.10%
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```
AAA47607 standard; cDNA; 2505 BP.
Human CARK (Cardiac related Ankyrin-Repeat Protein Kinase) cDNA.
WO200034330-A1.
 ABQ76480 standard; cDNA; 2531 BP.
S. cerevisiae BAX-associated cDNA fragment SEQ ID 385
 33
13
 2
11
19
 10
 14
39
 14
39
13
 12
25
7
 111
 AASSU426 standard; CDNA; 2495 BP.
DNA encoding novel human diagnostic protein #26230.
WO200175067-A2.
 Human secreted protein encoding cDNA SEQ ID NO 190. WO200277186-A2.
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indels:
 Conservative:
Mismatches:
Indel8:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Mismatches:
 Mismatches:
 ADC20175 standard; DNA; 2450 BP.
Human secreted protein coding sequence #114
WO200292787-A2.
 Indels:
 Indels:
 Indels:
 Indels:
 AAF44702 standard; cDNA; 2508 BP.
Novel protein kinase cDNA, SEQ ID NO: 83.
WO200073469-A2.
 Human polynucleotide SEQ ID NO 2137.
WO200157190-A2.
 BP.
 ABZ67070 standard; cDNA; 2450 BP
 Human MARK3-associated cDNA #60
US2003232771-A1.
 ADI29400 standard; cDNA; 2508
 MOLOCI-

03-OCT-

(HUMA-) HUMAN GENOME SCI INC.

ent Similarity: 40.91%

Local Similarity: 25.00%
 PA (MILL) MILLENNIUM PHARM INC.
Percent Similarity: 42.31%
Best Local Similarity: 38.46%
Query Match: 13.10%
 21-NOV-2002.
(HUMA-) HUMAN GENOME SCI INC.
 ID ADI29400 standard; curr, coll be Human MARK3-associated cDNA # US2003232771-A1.

PD 18-DEC-2003.

PA (ISIS-) ISIS PHARM INC.
Percent Similarity: 42.31%
Best Local Similarity: 38.46%
 (HUMA-) HUMAN GENOME SCI INC.
 42.31%
38.46%
13.10%
 46.55%
29.31%
13.10%
 40.91%
25.00%
13.10%
 unt Similarity: 40.91%
Local Similarity: 25.00%
Match: 13.10%
 46.67%
26.67%
13.10%
 PA (JANC) JANSSEN PHARM NV.
Percent Similarity: 46.55 Best Local Similarity: 29.31 Query Match: 1101
RESULT 1101
 ID ABZ67070 standard; of Human secreted prote PN WO2002/7186-A2.
PD 03-OCT-2002.
PA (HUMA-) HUMAN GENOME Percent Similarity: 2 Best Local Similarity: 2
 Percent Similarity:
Best Local Similarity:
 PA (HYSE-) HYSEQ INC.
Percent Similarity:
Best Local Similarity:
 PA (SUGE-) SUGEN INC. Percent Similarity:
 Local Similarity:
 ID ADC20175 standar BW Human secreted pW WO20029787-A2.
PD 21-NOV-2002.
PA (HUMA-) HUMAN GE Percent Similarity;
Best Local Similarity
 WO200264766-A2.
WO200277013-A2.
 15-JUN-2000.
 07-DEC-2000
 22-AUG-2002
 11-OCT-2001
 Query Match:
RESULT 1095
 Query Match:
RESULT 1096
 Match:
 Best Local S
Query Match:
RESULT 1094
 Match:
 Match:
 Query Match
RESULT 1098
 Query Match:
RESULT 1100
 Query Match
RESULT 1099
 Query Match
RESULT 1097
 Best
 PN
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SEQ ID NO 40489
 15
34
16
 15
34
16
 15
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16
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10
31
 19
25
25
 11 13
 12
43
21
 cDNA; 2788 BP.
human diagnostic protein #16524
 human diagnostic protein #19588
 ADOS7041 standard; cDNA; 3001 BP.
Human cardiac ankyrin repeat kinase, CARK, cDNA.
WO20040414164-A2.
27-MAY-2004.
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Conservative:
 Conservative:
 Conservative:
Mismatches:
 Conservative:
Mismatches:
 Conservative:
 Drosophila melanogaster genomic polynucleotide WO200171042-A2.
 Mismatches:
Indels:
 Mismatches:
Indels:
 Mismatches:
 Mismatches:
 Indels:
 Indels:
 Indels:
 Indels:
 Human polynucleotide SEQ ID NO 2139. W0200157190-A2.
 Human cDNA sequence SEQ ID NO:17956.
 Query Match: 13.10* In
RESULT 1102
DD AAK22609 standard; cDNA; 2553 BP.
DE Human polymucleotide SEQ ID NO 2138.
PN WO200157190-A2.
 Prokaryotic essential gene #33968.
WO200277183-A2.
 CDNA, 2805 BP
 RESULT 1108
ID ABL29672 standard; DNA; 2930 BP.
 AAH18100 standard; cDNA; 2626
 DNA; 2574
 PN W0200277183-AZ.
PD 03-OCT-2002.
PA (ELIT-) ELITRA PHARM INC.
Percent Similarity: 46.75%
Best Local Similarity: 33.77%
 44.44%
27.78%
13.10%
 44.44%
27.78%
13.10%
 43.18%
21.59%
13.10%
 40.22%
30.43%
13.10%
 44.44%
27.78%
13.10%
 42.31%
38.46%
 PA (PADA / FE 36.00$
Percent Similarity: 24.00$
Pest Local Similarity: 24.00$
Onerv Match: 13.10$
 13.10%
 (HELI-) HELIX RES INST.
 DE DNA encoding novel h
PN W0200175067-A2.
PD 11-OCT-2001.
PA (HYSE) HYSEQ INC.
Percent Similarity: 4
Best Local Similarity: 3
 DE Human polynucleotide
PN W0200157190-A2.
PD 09-AUG-2001.
PA (HYSE-) HYSEO INC.
Percent Similarity: 4
Best Local Similarity: 2
PD 09-AUG-2001.

PA (HYSE-) HYSEQ INC.

Percent Similarity:

Best Local Similarity:
 RESULT 1103
ID AAK52610 standard;
 DNA encoding novel WO200175067-A2.
 PD 11-OCT-2001.
PA (HYSE-) HYSEQ INC.
Percent Similarity:
Best Local Similarity:
 (PEKE) PE CORP NY.
 ACA52311 standard;
 AAS83784 standard;
 AAS80720 standard;
 PA (HYSE-) HYSEQ INC. Percent Similarity:
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 EP1074617-A2.
07-FEB-2001.
 27-SEP-2001
 09-AUG-2001
 Query Match:
RESULT 1105
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 RESULT 1106
 RESULT 1104
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Query Match: RESULT 1118 ID ABL11137 standa: DE Drosophila melan	90	it di	PN WO200157182-A2. PD 09-AUG-2001. PA (HUMA-) HUMAN GJ Percent Similarity:	it it	PN WO200157182-A2. PD 09-AUG-2001. PA (HUMA-) HUMAN GJ Percent Similarity: Rest Local Similarity:	Query Match: RESULT 1121 ID ACC72050 standa: DE BCU0715C gene #	FN WOZU03029421-AZ PD 10-APR-2003. PA (ORIG-) ORIGENE PATCENT Similarity.	Best Local Similarity Query Match: RESULT 1122 ID AAS79155 standar	DE DNA ENCOAING PN WOZOOI75067-AZ. PD 11-0CT-2001. PA (HYSE-) HYSEQ IN	st st sury	PN WO2003029421-A2 PD 10-APR-2003. PA (ORIG-) ORIGENE Percent Similarity:	Best Local Similarity Query Match: RESULT 1124 ID ACC72048 standar	<u>م</u> ر	Best Local Similarity Query Match: RESULT 1125 ID ADDRISOS Standar	DE Leukaemia-relate PN W0200303443-A2 PD 15-MAY-2003. PA (DEKR-) DEUT KRE PA (UYLU-) UNIV LUI PA (HAFE/) HAFERLAC PA (SCHO/) SCHOCH (
Conservative: 2 Mismatches: 11 Indels: 19	. BP. for cardiac therapeutic preparation.	INC. Conservative: 2 Mismatches: 11 Indels: 19	BP. Kyrin-Repeat Protein Kinase) cDNA.	Conservative: 2 Mismatches: 11 Indels: 19	BP. repeat protein kinase (CARK) cDNA.	Conservative: 2 Mismatches: 11 Indels: 19	P. ent SEQ ID NO: 49559.	Conservative: 9 Mismatches: 24 Indels: 5	p. c polynucleotide SEQ ID NO 40567.	Conservative: 12 Mismatches: 43 Indels: 21	3165 BP. diagnostic protein #26226.	Conservative: 12 Mismatches: 25 Indels: 7	P. ent SEQ ID NO: 54497.	Conservative: 9 Mismatches: 24 Indels: 5	p. promoter P30.  DRP. Conservative: 9 Mismatches: 17
PA (SEQU-) SEQUENOM INC. Percent Similarity: 42.31% Best Local Similarity: 38.46% Query Match: 13.10%	69 standard; DNA; 3001 DC51086 for screening 050894-A2.	ice St.	6 standard; ARK (Cardia: 4330-A1.	PD 15-JUN-2000. PA (MILL-) MILLENNIUM PHARM INC. Percent Similarity: 42.31% Best Local Similarity: 38.46% Query Match: 13.10%	RESULT 1112  ID ACC48580 standard; CDNA; 3025 BP.  DB Human cardiac-related ankyrin-repeat protein kinase PN WC2003020912-A2. PD 11-MAR003-	PA (MILL) MILLENNIUM PHARM INC. Percent Similarity: 42.31% Best Local Similarity: 38.46% Query Match: 13.10%	AESCLI 1113 ID AAC46278 standard; DNA; 3108 BP. DE Arabidopsis thaliana DNA fragment PN EP1033405-A2.	iry at	AESOLI 1114 ID ABL29698 standard; DNA; 3131 BP. DB Drosophila melanogaster genomic polynucleotide PN W0200171042-A2.	PD 27-SEP-2001. PA (FEKE) PE CORD NY. Percent Similarity: 36.00% Best Local Similarity: 24.00% Query Match: 13.10%	cDNA; human	PA (HYSE-) HYSEQ INC. Percent Similarity: 46.67% Best Local Similarity: 26.67% Query Match: 13.10%	ID AAC47620 standard; DNA; 3187 BP. DB Arabidopsis thaliana DNA fragment PN EP10334055-A2. PD 06-SEP-2000.	Percent Similarity: 47.27% Best Local Similarity: 30.91% Query Match: 13.10% RRSUT 1117	ID ADD93580 standard; DNA; 3228 BP.  DE Arabidopsis pathogen-inducible promoter P3 PN WO2003083042-A2.  PD 09-OCT-2003.  PA (DNAP ) DNA PLANT TECHNOLOGY CORP.  Percent Similarity: 49.06% Conserva  Best Local Similarity: 32.08% Mismatch

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ard; DNA; 3372 BP.
aematopoietic antigen genomic sequence SEQ ID NO:33977.
 ard; DNA; 3371 BP.
.aematopoietic antigen genomic sequence SEQ ID NO:33976.
 anogaster expressed polynucleotide SEQ ID NO 27893.
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 ard; cDNA; 3549 BP.
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 ard; DNA; 3664 BP.
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Conservative: 6 Mismatches: 18 Indels: 8	olynucleotide SEQ ID NO 22960.	Conservative: 14 Mismatches: 17 Indels: 17	antigen DNA SEQ ID NO: 8570.	Conservative: 18 Mismatches: 29 Indels: 5	DNA fragment SEQ 1D NO: 3098.	Conservative: 18 Mismatches: 29 Indels: 5	Q IN NO:29.	Conservative: 18 Mismatches: 29 Indels: 5	antigen DNA SEQ ID NO: 8571.	Conservative: 18 Mismatches: 29 Indels: 5	antigen DNA SEQ ID NO: 8572.	Conservative: 18 Mismatches: 29 Indels: 5	DNA fragment SEQ ID NO: 3100.	Conservative: 18 Mismatches: 29 Indels: 5	DNA fragment SEQ ID NO: 3099.
94 623 108 8	MESOLI 1120 DE ABL23829 standard; DNA; 3795 BP. DE Drosophila melanogaster genomic polynucleotide PN W0200171042-A2. PD 27-SEP-2001.	PA (PEKE) PE CORP NY.  Percent Similarity: 48.48%  Best Local Similarity: 27.27%  Query Match: 13.10%	KESULI 112/ ID AAL05882 standard, DNA; 3898 BP. DE Human reproductive system related PN W0200155320-A2.	PA CZ-AUG-ZOUJ. PA (HUMA-) HUMAH GENOME SCI INC. Percent Similarity: 51.43% Best Local Similarity: 25.71% Query Match:	RESULT 1128  ID ABL9846 standard; DNA; 3898 BP.  DE Human testicular antigen encoding DNA fragment PN W0200155317-A2.  PD 02-AUG-2001.	PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 51.43\$ Best Local Similarity: 25.71\$ Query Match: 13.10\$	ID AD46959 standard; DNA; 3898 BP. DE Human protein-encoding gene 5, SE PN WO200272763-A2.	PD 19-SEP-2002. PA (HUMA-) HUMA GENOME SCI INC. Percent Similarity: 51.43 Conservati Best Local Similarity: 25.71% Mismatchee Query Match: 13.10% Indels:	AALOS Standard; DNA; 3899 BP. DE Human reproductive system related PN WO200155320-A2.	it it	ID AALOS884 standard, DNA; 3899 BP. DE Human reproductive system related PD W0200155320-A2. PD 02-AUG-2001.	iry it	AESOLI 1135  ESOLI 1135  ESOLI 1135  DE Human testicular antigen encoding DNA fragment PN WQ200153317-A2. PD 02-4162-2001	PA (HUMA-) HUMAN GENOME SCI INC. Percent Similarity: 51.43% Best Local Similarity: 25.71% Query Match: 13.10%	DE Human testicular antigen encoding DNA fragment PN WO200155317-A2. PD 02-AUG-2001. PD (HUMA-) HUMAN GENOME SCI INC.

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nsmembrane protein (PRO) cDNA #175.
 cDNA; 3934 BP.
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 cDNA; 3933 BP.
737) cDNA sequence SEQ ID NO:314.
 DNA; 3934 BP. the identification of proteins.
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 cDNA; 3934 BP.
PRO polypeptide sequence #175.
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DE Bullan Secreted polypeptide FRO1431-encouing PN US2003040603-A1.	iypeptide rkoi	.431-encoding conA,	, SEQ ID NO:349.	DE Human secreted PN US2003040054-A
Percent Similarity:	43.96%	Conservative:	17	Percent Similarity:
Best Local Similarity: Query Match:	25.27% 13.10%	Mismatches: Indels:	33 12	Best Local Similari Query Match:
Ĕ	cDNA; 3934 E			RESULT 1152 ID ACA88757 stand
DE Human secreted polypeptide PRO1431-encoding PN US2003040064-A1.	lypeptide PRO1	.431-encoding cDNA,	, SEQ ID NO:349.	
FD Z7-FEB-2003. Percent Similarity:	43.96%	Conservative:	17	PD 20-FEB-2003. Percent Similarity:
Query Match:	13.10%	Indels:	120	Query Match:
ID ACD21957 standard; cDNA; 3934 BP.	cDNA; 3934 E	ל זארן נ	777	ID ACA70199 stand
	insuce more and	e di di		DE MANAGESTA PRI US2003036134-A. PD 20-FEB-2003.
Percent Similarity: Best Local Similarity:	43.96% 25.27%	Conservative: Mismatches:	17 39	Percent Similarity: Best Local Similari
Query Match: RESILT 1145	13.10%	Indels:	12	Query Match: RESTET 1154
ID ACF13122 standard, cDNA, 3934 BP.  E Human secreted polypeptide PRO1431-encoding	cDNA; 3934 BP Preptide PR014	P. .431-encoding cDNA,	, SEQ ID NO:349.	ID ACTIVATION STANDS DE NOVEL human sec
	;			PD 30-JAN-2003.
Percent Similarity: Best Local Similarity:	43.96% 25.27%	Conservative: Mismatches:	17 39	Percent Similarity: Best Local Similari
Query Match: RESULT 1146	13.10%	Indels:	12	Query Match: RESULT 1155
ID ACD25225 standard; DE Human secreted/tran	; cDNA; 3934 BP	3P.	#175.	ID ACC74336 stand
PN US2003044925-A1. PD 06-MAR-2003	•			PN US2003027275-A
i ce	43.96%	Conservative:	17	n G
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4 stand	CDNA; 3934 E	,		700
DE Human secreted pol PN US2003054474-A1.	polypeptide PRO1431-encoding 1.	.431-encoding cDNA,	, SEQ ID NO:349.	DE Human secreted PN US2003027324-A
20-MAR-2003.	CMT			PD 06-FEB-2003.
cent Similarity:	-	Conservative:	17	Percent Similarity: Best Local Similari
Best Local Similarity: Onerv Match:	25.27%	Mismatches: Indels:	98	Query Match: RESILT 1157
ĭğ.			<b>3</b>	ID ACD25532 stand
		mbrane protein	PRO1431 CDNA.	
FN USZUU3U3ZII4-AI. PD 13-FEB-2003.				PD 20-FEB-2003. PA (GETH ) GENENT)
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rce at	43.96%	Conservative: Mismatches:	17	n Ce
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ID ACD18316 standard;	cDNA; 3934 E	4 NG 2	#175	ID ACC88296 stand
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12	#175.	17 39 12	PRO1431 cDNA.	17 39 12	#175.	17 39 12	PRO1431 cDNA.	17 39 12	SEQ ID NO:349.	17 339 12	175.	117 339 12	PRO1431 CDNA.	17 339 12	175.	17 39 12	SEQ ID NO:349.	17 39
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13.10* 17 1160 T 1160 ACD21650 standard; CDNA; 3934 BP. Human secreted/transmembrane protein (PRO)
Conservative Mismatches: Indels:
T 1161 PCD1817 standard; cDNA; 3934 BP. Human secreted/transmembrane protein (PRO) US2003044916-A1.
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Jr 1164 2009868 standard; cDNA; 3934 BP. Human secreted/transmembrane protein (PRO) US2003036128-A1.
Conservative: Mismatches: Indels:
Jr 1165 ACC88603 standard; cDNA; 3934 BP. Human secreted polypeptide PRO1431-encoding VS20030027266-Al.
Conservative Mismatches: Indels:
ALTISO ALTISO Human secreted/transmembrane protein (PRO) US2003054483-Al.
Conservative Mismatches: Indels:
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Query Match: RESULT 1169 ID ACA97394 standard; c DE Novel human secreted	13.10% Inde CDNA; 3934 BP. ed and transmembrane	ls: protein	12 PRO1431 CDNA.
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1 ~1	3.10%	Indels:	12
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ery Match: SULT 1171	) 0 0	Indels:	12
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<i>,</i> ,	<b>*</b> 96.	Conservative:	17
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p. 131-encoding	, SEQ ID NO:349.	ID ACA95323 standard DE Novel human secret PN US2003032119-A1.
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		1D ACD12746 standard; cDNA; DE Human secreted/transmemb PN US2003016125-A1.
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(PRO) cDNA	#175.	IDS ACTIONS STANDARY DE Human secreted pr PN US2003040068-Al.
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ein (PRO) cDNA	#175.	TESOLI 1195  ID ABX76920 standard  DE Human PRO polynuc PN US2003027280-A1.
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protein	RO1431 CDNA.	ID ACA73252 standard DE Novel human secrei PN US20032300-AI.
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protein	RO1431 CDNA.	TESOLI 1134 ID ACA68795 standard; CDNA; DE NOVEL human secreted and PN US2003036136-A1.
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4	BB. Conservative: Mismatches: Indels: BP. Smembrane protein Conservative: Mismatches: Indels:	g cDNA, SEQ ID NO:34 es: 39 g cDNA, SEQ ID NO:34 tive: 17 es: 39 cDNA #175. tive: 17 es: 39 tein PRO1431 cDNA. tive: 17 es: 39 tein PRO1431 cDNA. tive: 17 es: 39 conA #175. tive: 17 es: 39 conA #175.

ttch:  (6) 13 similar  (6) 13 stan  (6) 13 stan  (6) 13 stan  (6) 13 stan  (7) 18 stan  (8) 18 stan  (9) 18 s	.27% Mismatches: 39 .10% Indels: 12 NA; 3934 BP. and transmembrane protein PRO1431 cDNA.	5.96% Conservative: 17 5.27% Mismatches: 39 5.10% Indels: 12	NA; 3934 BP. ptide PRO1431-encoding cDNA, SEQ ID NO:349.	.27% Conservative: 17 .27% Mismacches: 39 .10% Indels: 12	NA; 3934 BP. ptide PRO1431-encoding cDNA, SEQ ID NO:349.	.27% Conservative: 17 .27% Mismacches: 39 .10% Indels: 12	NA, 3934 BP. embrane protein (PRO) cDNA #175.	96% Conservative: 17 .27% Mismatches: 39 .10% Indels: 12	NA; 3934 BP. ptide PR01431-encoding cDNA, SEQ ID NO:349.	.06% Conservative: 17 .27% Mismatches: 39 .10% Indels: 12	NA; 3934 BP. ide #175.	.27% Conservative: 17 .27% Mismatches: 39 .10% Indels: 12	NA; 3934 BP. and transmembrane protein PRO1431 cDNA.	.96% Conservative: 17 .27% Mismatches: 39 .10% Indels: 12	DNA; 3934 BP. and transmembrane protein PRO1431 cDNA.	.96% Conservative: 17 .27% Mismatches: 39 .10% Indels: 12	NA; 3934 BP. RO polypeptide #175.
	<pre>Y: 25. 13. rd; cDN reted a</pre>	003. GENENTECH INC arity: 4: milarity: 29	standard; cl creted polype 7263-A1.	003. arity: 43. milarity: 25. 13.	1189 CC90138 standard; cDNA, uman secreted polypept; S2003027271-A1.	5-FEB-2003. : Similarity: 43.9 ocal Similarity: 25.2 Aatch: 13.1	1150 2012746 standard; cDNA; 3934 Jman secreted/transmembrane 32003036125-A1.		1131 2F19976 standard; cDNA; 1man secreted polypeptio 32003040068-Al.		3X76920 standard; cDNA; lman PRO polynucleotide 52003027280-A1.		8	. n in in	5 standard; c uman secreted 36136-Al.	03. rity: 43. ilarity: 25.	standard; cDN ding human PR 138-A1.

Mismatches:

est Local Similarity: 25.27% uery Match: ESULT 1196	74 94 94	Mismatches: Indels:	39 12	Best Local Similarity: 25.27% Query Match: RESULT 1205
ACA70506 standard; cDNA; 393 Human secreted/transmembrane US2003032109-A1.	; 3934 BP. brane protein	ein (PRO) cDNA	#175.	ID ACA96366 standard; cDNA; 39. DE Human PRO polynucleotide #1. PN US2003017540-A1.
PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	9	Conservative: Mismatches: Indels:	17 39 12	ar de
KESULI 119/ DD ACD14692 standard; cDNA; DE Human PRO polynucleotide PN US2003040066-Al.	; 3934 BP. e #175.			RESULT 1206  ID ACA65140 standard; cDNA; 39: DE Human PRO polynucleotide #1: PN US2003032106-A1.
PD 27-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 110%		Conservative: Mismatches: Indels:	17 39 12	PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%
ACA68364 standard; CDNA; DE Novel human secreted and PN US2003032104-A1.	; 3934 BP. d transmembrane	protein	PRO1431 CDNA.	70.5
PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%		Conservative: Mismatches: Indels:	17 39 12	PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%
KESOLI 1199 ID ABX98829 standard; CDNA; DE Novel human secreted and PN US200305157-A1.	; 3934 BP. d transmembrane	protein	PRO1431 cDNA.	RESULT 1208 ID ACA74278 standard; cDNA; 39: DE Novel human secreted and tri PN US2003032131-A1.
PD ZU-FEB-ZU03.  Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	0888 088	Conservative: Mismatches: Indels:	17 39 12	الم الم الم
KESULT 1200 DE ACCESSOR Standard; CDNA; 39 DE Human secreted polypeptide PN US2003032120-A1.	; 3934 BP. ide PRO1431-	1-encoding cDNA,	, SEQ ID NO:349.	RESULT 1209 ID ACA96673 standard; cDNA; 39; DE Human PRO polynucleotide #1' PN US2003032133-Al.
PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	0 7 6 % % %	Conservative: Mismatches: Indels:	17 39 12	PD 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query_Match: 13.10%
ACA95630 standard; cDNA; Novel human secreted and US2003036155-A1.	; 3934 BP. d transmembrane	protein	PRO1431 cDNA.	RESULT 1210 ID ACD10779 standard; cDNA; 39: DE cDNA encoding human PRO pol, PN US2003032107-A1.
Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	7* 7*	Conservative: Mismatches: Indels:	17 39 12	FU 13-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%
1202 ACD04548 standard; cDNA; Novel human secreted and US2003022296-A1.	; 3934 BP. d transmembrane	protein	PRO1431 CDNA.	RESULT 1211 ID ACC91475 standard; cDNA; 39: DE Human secreted polypeptide in US2003031139-A1.
Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10% BESTITT: 1203		Conservative: Mismatches: Indels:	17 39 12	Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%
87989 standard; lan secreted pol 1003027281-A1. FEB-2003.	, 3934 BP. ide PRO1431	3934 BP. le PRO1431-encoding cDNA,	, SEQ ID NO:349.	IESULI 1212 ID ACD02810 standard; CDNA; 39: DE CDNA encoding human PRO poly PP US2003022301-A1. PD 30-7AN-2003
PA (GETH ) GENENTECH INC. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%		Conservative: Mismatches: Indels:	17 339 12	ccent Similarity: 43.96% st Local Similarity: 25.27% ery Match: 13.10% sULT 1213
51 standa secreted 040058-A1	ırd; cDNA; 3934 BP. polypeptide PR01431-encoding	1-encoding cDNA		1D ACC8/3/5 Standard; CDNA; 39: DE Human secreted polypeptide F PN US2003036165-A1. PD 20-FEB-2003. PA (GETH ) GENENTECH INC.
ercent Similarity: 43.96%	**	Conservative:	17	rcent Similarity:

а С	est botal similarity;	#/7:C7	Transcorter:	20.0
וצמגו	RESULT 1205	9	מ	
- Ω	standard; O polynucl	cDNA; 3934 BP. eotide #175.		
ሷ ሷ	7540-A1.			
4 Д	ercent Similarity:	3.96%	Conservative:	
Д	Best Local Similarity:	25.27%	Mismatches:	39
OΥP	Query Match: RESIII.T 1206	13.10%	Indels:	
H	D ACA65140 standard;	DNA; 393		
ΩĤ	E Human PRO polynucle	otide #1		
<b>1</b> 4 P4	N USZU0303Z106-A1. D 13-FEB-2003.			
Д	ercent Similarity:	13.96\$	Conservative:	17
ш	est Local Similarity:	25.27%	Mismatches:	39
ZI EZ	Query Match: RESULT 1207	3.10%	Indels:	12
н	D ACA73866 standard;	DNA; 3934 BP.		
ΩΩ	DE Human secreted/trans	membrane prot	ein (PRO) cDNA	#175.
ı Di	D 13-FEB-2003.			
Д	ercent Similarity:	3.96%	Conservative:	17
шĊ	est Local Similarity:	5.27%	Mismatches:	39
X KZ	ESULT 1208			71
ні	D ACA74278 standard;	CDNA; 3934 BP.		
O O	E Novel human secrete	and t	protein	PRO1431 cDNA.
ı Di	N USZUUSUSZISI-AI: D 13-FEB-2003.			
Δi	ercent Similarity:	3.968	Conservative:	
ρά ·	est Local Similarity:	5.27%	Mismatches:	39
Or p	uery Match: pent 1200	3.10%	Indels:	
ΥÄ	ESULI 1203 D ACA96673 Standard:	F98 : AND		
Ä	E Human PRO polynucle	tide #175.		
A i	N US2003032103-A1.			
ם, כ	D 13-FEB-2003.			
שיע	ercent Similarity: est Local Similarity:	3.26*	Conservative:	17
άÓ	est botal similarity: merv Match:	3.10%	מ	
i izd	ESULT 1210		100	
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O 0	E CDNA encoding humar	PRO	le #175.	
i A	0820030310/-AL.			
ñ	ercent Similarity:	3.96\$	Conservative:	17
Ē	est Local Similarity:	5.27%		39
Ö	lery Match:	3.10%	Indels:	12
¥ FI	ESULI 1211 D. ACC91475 standard:	DNA: 30		
Ö	E Human secreted poly	eptide	-encoding cDNA,	SEQ ID NO:349.
<u>a</u> a	N US2003032139-A1.			
ri U	o is-reb-2003. Proent Similarity.	3 968	Conservative.	7.1
ищ	est Local Similarity:	5.278	)	39
δį	lery Match:	3.10%	Indels:	12
24 F	ESULT 1212	ים גנסני גאות		
DE	s cDNA encoding human	PRO polypeptide	e #175.	
NG	US2003022301-A1.	3-37-3	! :	
ā	۲.	890	4	
m	Best Local Similarity:	5.278	Mismatches:	39
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áa:	ACC87375 standard; cDNA;	3934 BP.		
D E		le PRO1431	-encoding cDNA,	SEQ ID NO:349.
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PA Pe	2	INC.	Conservative.	17
•	rcent similarity:	43.304	COlliser valiate.	7.

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ery Match: SULT 1223 ACA74946 standard;	mar	Percent Similarity: 45,703 Best Local Similarity: 25,278 Query Match: 13,108	RESOLI 1224 ID TAMAN PRO POLYMICLEOTIDE	PN US2003032128-A1. PD 13-FEB-2003.	Percent Similarity: 43.96* Best Local Similarity: 25.27*	ery Matcn: SULT 1225 ACA71461 standard:	tu; tran		Best Local Similarity: 25.27% Query Match: 13.10%	Ë	creted poly 2122-Al.	ပ္ပိ	Best Local Similarity: 25.27% Ouerv Match: 13.10%	SULT 1227			Percent Similarity: 43.203 Best Local Similarity: 25.27\$	SULT 1228		•	Percent Similarity: 43.96%  Best Local Similarity: 25.27%	13	ID ACD16578 standard; cDNA;	PN US2003017543-A1. PD 23-JAN-2003.	rge +		ID ACD15657 standard; cDNA; 3934 DE Human secreted/transmembrane		Percent Similarity: 43.964  Best Local Similarity: 25.278  Onerw Match:	•
39 12	SEQ ID NO:349.	17	7		F-6	2	7. 5.		69	[2]			68		PRO1431 CDNA		21	12		PRO1431 cDNA.	2.1	399		#175.	-	39 12	1	#175.	1.7 3.0	12
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25.27% 13.10%	cDNA; 3934 BP. ypeptide PRO1431	43.96%	13.10% CDNA: 393	sotide #175.		25.27% 13.10%	CDNA; 3934 BP.	allend talle	43.96% 25.27%	13.10%	cDNA; 3934 BP.		43.96%	) (M	CDNA; 3934 BP.		3.96*	25.27* 13.10*	CDNA: 3934 BP.	ed and transmembrane	<b>.</b>	25.27%	14. 2024 BD	u, coma; 333% pr. ransmembrane protein	* * *	13.10%	DNA; 3934 BP.	orane prot	43.968	13.10%
Best Local Similarity: Query Match: RESHLT 1214	ID ACC85959 standard, cDNA; 3934 BP. DE Human secreted polypeptide PRO1431 PN US2003027262-A1.	PD 06-FEB-2003. Percent Similarity: Best Local Similarity:	Query Match: RESULT 1215 ID ACA65447 standard;	Human PRO polynuch US2003032110-A1.	PD 13-FEB-2003. Percent Similarity:	Best Local Similarity: Query Match:	RESULT 1216 ID ACA94264 standard; DE Himan Secreted/tran		ž č	Query Match:	08 standard; PRO polynuci	US2003036145-A1.	i ce	Query Match:	RESULT 1218 ID ACA91510 standard; CI DE Nextel himan secreted	PN US2003036154-A1.	PD 20-FEB-2003. Percent Similarity:	Best Local Similarity: Query Match:	RESULT 1219 ID ACA90724 standard:		ŭ	Best Local Similarity:	RESULT 1220	DE Human secreted/tra PN 1152003044931-A1.	06-MAR-2003	Fercenc Similarity: Best Local Similarity: Onerv Match:	RESULT 1221 ID ACD17432 standard:	Human secreted/t US2003036150-A1	PD 20-FEB-2003. Percent Similarity:	

		31 cDNA.		Q ID NO:349.			31 cDNA.												0 TD NO:349				Q ID NO:349.		0 4 5. OM OT O
	17 39 12	PRO1431	17 39 12	, SEQ		39	12 PRO1431		17	12	#175	1 1	17	12	#175	) 4	17	12	Q.		17	12	, SEQ	39	1 U
	Conservative: Mismatches: Indels:	protein	Conservative: Mismatches: Indels:	1-encoding cDNA,		conservative: Mismatches: Indele:	protein		Conservative:	Mismatches: Indels:	(PRO) CDNA		Conservative:	Indels:	אַאַרוּהַ (PRO) תוֹפּ		Conservative:	Mismatches: Indels:	1-encoding only		Conservative:	Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indela:	
cDNA; 3934 BP. eotide #175.	43.96% 25.27% 13.10%	cDNA; 3934 BP. ed and transmembrane	43.96% 25.27% 13.10%	CDNA; 3934 BP. Ypeptide PR0143	INC.	43.96% 25.27% 10.10%	cDNA; 3934 BP.	Ç	43.96%	25.2/\$ 13.10%	cDNA; 3934 BP.		43.96%	13.10%	CDNA; 3934 BP.		43.96%	25.27% 13.10%	CDNA; 3934 BP.		43.96%	13.10%	cDNA; 3934 BP. ypeptide PR0143	INC. 43.96% 25.27% 13.10%	CDNA; 3934 BP.
ID ACA97701 standard; cDNA; DE Human PRO polynucleotide PN US2003032115-A1.	PD 13-FBB-2003. Percent Similarity: Best Local Similarity: Query Match:	ACA99150 standard; CI Novel human secreted IIS2003032140-Al	PD 33-FEB-2003. Percent Similarity: Best Local Similarity: Query Match:	Jur 1234 ACC91782 standard; cDNA; 3934 BP. Human secreted TRSP03040076.21	575	imilarity:	RESULT 1235 ID ACD11193 standard; cl DE Novel human secreted	US2003008352-A1. 09-JAN-2003.	C	sımılarıty: :	KESULT 1236 ID ACD15043 standard; CDNA; 3934 BP. DE Human secreted/transmembrane protein	US2003044922-A1.	Forcent Similarity: Rest Local Similarity:		DE Human secreted/transmembrane protein	US2003032118-A1.	ty:	Best Local Similarity: Query Match:	RESOLI 1236 ID ACC95936 standard; CDNA; 3934 BP. DF. Himms secreted nolvmentide PRO1431-encoding	US2003036135-A1. 20-FEB-2003.	larity:	: Local Similarity: cy Match: ILT 1239	<pre>1D ACF16499 standard; cDNA; 3934 BP. DE Human secreted polypeptide PR01431-encoding PN US2003054455-A1.</pre>	003. GENENTECH arity: milarity:	7 standard

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ACF78088 standard; cDNA; 3934 BP.
ACF78088 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003554479-Al.
20-MAR-2003.
(GETH) GENEWTECH INC.
(GETH) GENEWTECH INC.
xcent Similarity: 43.96$ Mismatches: 39
xxv. March: 13.10$ Indels: 12
 ULT 1241
ACRO20243 standard; CDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349.
US2003049743-A1.
13.0ARA-2003.
(GETH) GENENTECH INC.
cent Similarity: 43.96* Mismatches: 17
Local Similarity: 13.10* Indels: 12
ULT 1242
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003068743-A1. (GETH) GENENTECH INC. (GETH) GENENTECH INC. Conservative: 17 t Local Similarity: 25.27% Mismatches: 39 Indels: 13.10% Indels: 12
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003068752-A1.
 ID NO:349.
 CDNA, SEQ
 ry Match: 13.10% Indels: 12
ULT 1245
ACD46793 standard; cDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
US2003068685-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175.
US2003068725-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175.
US2003068682-A1.
10-APR-2003.
(GETH) GENENTECH INC.
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Mismatches:
 ACF21511 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding
US2003049769-A1.
 Indels:
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 Jr 1247
ACF28323 standard, CDNA, 3934 BP.
 MT 1246
ACD49556 standard; cDNA; 3934 BP.
 CDNA; 3934 BP
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cent Similarity: 43.96*
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ecreted po 40071-A1. 2003.	PA (GETH ) GENENTECH INC.  Percent Similarity: 43.96\$ Conservative: 17  Best Local Similarity: 25.27\$ Mismatches: 39  Query Match: 13.10\$ Indels: 12	9 standard, ecreted/tra 54471-A1. 2003.	Percent Similarity: 43.96\$ Conservative: 17 Percent Similarity: 25.27\$ Mismatches: 39 Query Match: 13.10\$ Indels: 12	RESULT 1259 ID ACF18827 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. PN US2003064452-A1. PD 03-APR-2003.	9,37,52	4 standard ecreted po 68705-A1. 2003.	9 tr g	RESOLI 1201 ID ACT78395 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. PN US2003054473-A1.	PD 20-MRk-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 43.96% Conservative: 17  Percent Similarity: 25.27% Mismatches: 39  Court. March Similarity: 13.10% Tadals: 13	Type Reconstruction 1930 BP. ACF51994 standard; cDNA; 3934 BP. Human secreted polypeptide PRO1431-encoding cDNA,	PN 03-ARR-2003. PD 03-ARR-2003. Percent Similarity: 43.96* Conservative: 17 Best Local Similarity: 25.27* Mismatches: 39	DNA; 3934 BP. eptide PRO1431-encoding cDNA,	PD 10-APP-2003. PA (GETH ) GRENTECH INC. PA (GETH ) GRENTRECH 10.96\$  Percent Similarity: 25.27\$  Best Local Similarity: 25.27\$	Indels: 1934 BP. PRO1431-encoding cDNA,	US_ZUUJUBS/ZZ_AI. 10.ARF_2Z_003. (GETH ) GENENTECH INC. cent Similarity: 43.96% Conservative: 1	rd o
17 39 12		17 339 12		17 39 12	#175.	17 39 12	, SEQ ID NO:349.	17 39 12	#175.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	17 39 12
Conservative: Mismatches: Indels:	·	Conservative: Mismatches: Indels:	ide #175.	Conservative: Mismatches: Indels:	) cdna	Conservative: Mismatches: Indels:	334 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	cein (PRO) cDNA	Conservative: Mismatches: Indels:	334 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:
43.96% 25.27% 13.10%	cDNA; 3934 BP. eotide #175.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. n PRO polypeptide	INC. 43.96% 25.27% 13.10%	CDNA; 3934 BP. nsmembrane prot	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. ypeptide PRO143	43.96% 25.27% 13.10%	cDNA; 3934 BP. nsmembrane prot	43.96% 25.27% 13.10%		43.96% 25.27% 13.10%	cDNA; 3934 BP. Ypeptide PRO143	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP Ypeptide PRO143	43.96% 25.27% 13.10%
Percent Similarity: Best Local Similarity: Query Match:	ğ	GENENTECH ilarity: Similarity:	RESULT 1250 ID ACD99182 standard; OB CDNA encoding human PN US2003068755-A1.	PD 10-APR-2003. PA (GETH ) GENEWTECH Percent Similarity: Best Local Similarity: Query Match:	sur	-2003. ) GENENTECH ilarity: Similarity:	RESULT 1252  ID ACF48924 standard; cDNA; 35  DE Human secreted polypeptide PN US2003104539-Al.	FU US-UN-2005. Percent Similarity: Best Local Similarity: Query Match:	RESULT 1253 ID ACD09244 standard; cDNA; 3934 DE Human secreted/transmembrane PN US2003036131-A1.	rce st	KESULI 1254  ID ACF12007 standard, cDNA, 3: DE Human secreted polypoptide PN US2003040075-A1.	For 2, FEB-2003.  Percent Similarity: Best Local Similarity: Query Match:	71 standard secreted pol 054459-A1.	(GETH ) GENENTECH coent Similarity: St Local Similarity: ST Match:	rESULT 1256 ID ACF15885 standard; cDNA; 35 DE Human secreted polypeptide PN US2003044930-A1.	PD 06-MAR-2003.  Percent Similarity: Best Local Similarity: Query Match: RESULT 1257

HINC.  43.96% Conservative: 1 25.27% Mismatches: 3 13.10% Indels: 1 d; cDNA; 3934 BP. olypeptide PRO1431-encoding cDNA,	Percent Similarity: 43.96\$ Conservative: 17 Best Local Similarity: 25.27\$ Mismatches: 39 Query Match: 13.10\$ Indels: 12 RESULT 1275 ID ACE7402 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. PN US2003068699-Al. PD 10-APR-2003. PA (GETH) GENENTECH INC. Percent Similarity: 43.96\$ Mismatches: 39 Best Local Similarity: 25.27\$ Mismatches: 39	13.10% Indels: d; cDNA; 3934 BP. olypeptide PR01431-encoding cDNA, H INC. Conservative: 25.27% Mismatches: 13.10% Indels:	RESULT 1277  RESULT 1277  RESULT 1277  EXENCE 1277  DE Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349.  PU US2003073175-A1.  PD 17-APR-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 43.96% Mismatches: 39  Query Match: 13.10% Indels: 12	ACD89934 standard; CDNA; 3934 Hunan secreted/transmembrane p US2003068695-A1. 10-APR-2003. (GETH) GENENTECH INC. (GETH) GENENTECH Similarity: 25.27% 12. Macch: 13.10% ULT 1279 ctandard; CDNA: 3004	ictide #175. lectide #175. INC. 43.96% 25.27% 13.10% cDNA; 3934 B	rce st sury sur
17-APR-2003. (GETH ) GENENTECH rcent Similarity: st Local Similarity: ery Match: ery Match: ACF50459 standard Human secreted po. US2003104549-A1. 05-JUN-2003.	Conservative: Mismatches: Indels: 934 BP. ne protein (PRO) cDNA #	13.10% Misma 13.10% Indel d; cDNA; 3934 BP. olypeptide PRO1431-enco H INC. Conse : 25.27% Misma	Indels: 1 1934 BP. ane protein (PRO) cDNA #1 Conservative: 1 Mismatches: 3	13.10\$ Indels:   1   1   1   1   1   1   1   1   1	RESULT 1271 ID ACT23353 standard; CDNA; 3934 BP. ID ACT23353 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349. DE Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349. PN 19200306463-A1.  Conservative: 17  RESULT 1272 ID ACT40043 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349. PN 192003064463-A1.	rce st ery SUI

Best Local Similarity: 29 Query Match: 13	25.27% 13.10%	Mismatches: Indels:	39 12	PD 06-MA
1786 F76860 standard; man secreted poly 2003104548-A1.		334 BP. PRO1431-encoding cDNA,	, SEQ ID NO:349.	Best Local Query Matc RESULT 129 ID ACC97
05-JUN-2003. (GETH ) GENENTECH coent Similarity: it Local Similarity: rry Match:	INC. 43.96% 25.27% 13.10%	Conservative: Mismatches: Indels:	17 39 12	DB Human PN US200 PD 06 PD 06 PD 06 Percent Sill Best Local
5 standa ecreted 04542-A	ard; cDNA; 3934 BP. polypeptide PRO1431 1.	3934 BP. de PRO1431-encoding cDNA,	, SEQ 1D NO:349.	L L
003. arity: milarity:	43.96% 25.27% 13.10%	Conservative: Mismatches: Indels:	17 12 12	PN US2000 PD 27-MAI PA (GETH Percent Sir
SULT 1284 ACF50152 standard; Human secreted poly US2003104543-A1.	cDNA; 3934 BP. peptide PRO1431	encoding cDNA,	. SEQ ID NO:349.	it it
PD 05-JUN-2003. Percent Similarity: 43 Best Local Similarity: 25 Query Match: 13	43.96% 25.27% 13.10%	Conservative: Mismatches: Indels:	17 39 12	DE Human PN US2000 PD 03-API PA (GETH
resoni 1299 ID ACD09551 standard; cI DE Human secreted/transm	cDNA; 3934 BP.	(PRO) CDNA	#175.	Best Local
	43.96% 25.27% 13.10%	servative: matches:	17 39 12	RESULT 1299 ID ACF14, DE Human PN US200 PD 20-MAI
KESULI 1286 ID ACD08630 standard; cDNA; 3934 DE Human secreted/transmembrane DN US2003940061-A1.	cDNA; 3934 BP. smembrane protein	(PRO) cDNA	#175.	PA (GETH Percent Sir Best Local Onery Match
PD 27-FEB-2003.  Percent Similarity: 43 Best Local Similarity: 25 Boary Match: 13	43.96% 25.27% 13.10%	Conservative: Mismatches: Indels:	17 39 12	
ACF12344 standard; Human secreted poly US2003036130-A1.		134 BP. PRO1431-encoding cDNA,	SEQ ID NO:349.	PD 10-APE PA (GETH Percent Sir Best Local
at series		Conservative: Mismatches: Indels:	17 339 12	r F
2 standard; ecreted poly 54468-Al.	cDNA; 3934 BP. Peptide PRO1431	934 BP. PRO1431-encoding cDNA,	SEQ ID NO:349.	PN USZOUS PD 17-API PA (GETH Percent Sin
003. GENENTECH 1 arity: milarity:	968 278 108	Conservative: Mismatches: Indels:	17 39 12	Best Local Guery Match RESULT 1297 ID ACD48
ACD22571 standard; Human secreted/trar US2003054470-A1.	cDNA; 3934 BP. Ismembrane prote	CDNA	#175.	PN USZOU: PD 03-APF PA (GETH PACENT Sin
PD 20-MAR-2003.  A (GETH) GENENTECH INC Percent Similarity: 43 Best Local Similarity: 25 Ouery Match:	INC. 43.96% 25.27% 13.10%	Conservative: Mismatches: Indels:	17 39 12	Best Local Query Match RESULT 1296 ID ACD48
KESULI 1230 ID ACF15271 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding cDNA, PN US2003044917-A1.	ONA, 3934 BP. eptide PRO1431	encoding cDNA,	SEQ ID NO:349.	PN USZOU- PD 03-APF PA (GETH Percent Sin

: 17 39 12 18 SEO TD NO. 349	17 339 12	NA, SEQ ID NO:349.	ਜ਼ ਜ਼ੁਲ	12 DNA, SEQ ID NO:349.	17 12 12 12 12 12	NA, SEC 1D NO:349. 17 39 12	PRO1431 CDNA.	1 #1	17 339 12	, #175.
Conservative: Mismatches: Indels: 34 BP.	Conservati Mismatches Indels:	34 BP. PRO1431-encoding cDNA, Conservative: Mismatches:	a e	Indels: BP. 01431-encoding c	onservati ismatches ndels:	encoainy cu Conservative Aismatches: Indels:	34 BP. ansmembrane protein Conservative: Mismarches:	Indels: BP. protein (PRO) c	Conservative: Mismatches: Indels:	protein (PRO) cDNA
43.96% -Y: 25.27% 13.10% ard; cDNA; 39	43.96 <b>%</b> : 25.27 <b>%</b> : 13.10 <b>%</b>	a; cDNA; 39 olypeptide r INC. 43.96% c 25.27%	13.10% 13, CDNA; 35 51ypeptide 1 INC. 43.96%	13.10 cd; cDNA;	CH INC. 43.96% Y: 25.27% 13.10% rd, cDNA, 35	INC. 43.96% 25.27% 13.10%	; cDNA; 39 ted and tr INC. 43.96% 25.27%	13.10% ; cDNA; 393 ansmembrane	TECH INC. :: 43.96% ity: 25.27% 13.10%	cDNA, 3934 nsmembrane INC.
PD 06-MAR-2003. Percent Similarity: Best Local Similarity Query Match: RESULT 1291 ID ACC97366 stande DR Human secreted	PN US2003044929-A1 PD 06-MAR-2003. Percent Similarity: Best Local Similarit Query Match:		Query Match: RESULT 1293 ID ACF14043 standard; DE Human secreted pol PN US2003064465-A1. PD 03-APR-2003. PA (GFTH ) GENENTECH PErcent Similarity:	Query Match: RESULT 1294 ID ACF14350 standar DE Human secreted PN US2003054478-A1.	20-MA (GETH (GETH st Local st Local sry Matc SULT 129	DE MUMMAL SECTECHE JOH DE 10-APR-2003. PA (GETH) OBENUTECH PECCHT SIMILATICY: Best Local Similarity: Query Match:	ACCOUNT ACCESSED BY ACCOUNT AC	Query Match: RESULT 1297 ID ACD45872 stan DE Human secrete PN US2003064454- PD 03-APR-2003.	PA (GETH ) GENENTECH Percent Similarity: Best Local Similarity: Query Match: PERTIT 1998	1230 D48021 st man secre 200306446 -APR-2003 ETH ) GEN

Query Match: RESULT 1307 ID ACH1204	DE CONA en PN US20030. PD 13-MAR-	9 7 7 5		PAR (GETH ) Percent Simi Best Local S Query Match: RESULT 1309	ID ACD4074. DE Human so PN US20079. PD 13-FFEB-	£ # £ £	DE TUINGIL ST PN US20030 PD 20-MAR-, PA (GETH)	of the state of th	DE HUMAN SON SON SON SON SON SON SON SON SON SO	in the second	DE Human sv PN USC0030- PD 13-MAR PA (GETH)	g # # g	DE HUMBAIL SI	2 7 K E	DE Human sr PN US20030 PD 10-APR- PA (GETH )	Percent Simi: Best Local S: Query Match: RESULT 1315
2		2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	2 9 7	75.	7.66	v	2 9 7	#175.	7 6 7	SEQ ID NO:349.	2 9 7	SEQ ID NO:349.	7-29
Mismatches: 3 Indels: 1	e #175.	Conservative: 1 Mismatches: 3 Indels: 1	CDNA,	Lve:	CDNA,	Conservative: 1 Mismatches: 3 Indels: 1	in (PRO) cDNA #17	Conservative: 1		Conservative: 1 Mismatches: 3 Indels: 1	CDNA	Conservative: 1 Mismatches: 3 Indels: 1	cDNA,	Conservative: 1 Mismatches: 3 Indels: 1	CDNA,	Conservative: 1 Mismatches: 3
25.27% 13.10%	7752 standard; cDNA; 3934 BP. encoding human PRO polypeptide 73068724-A1.	INC. 43.96% 25.27% 13.10%	ırd; cDNA; 3934 BP. polypeptide PRO1431-encoding L.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. ypeptide PRO1431	INC. 43.96 <b>%</b> 25.27 <b>%</b> 13.10 <b>%</b>	cDNA; 3934 BP. nsmembrane prote	INC. 43.96% 25.27%	cDNA; 3934 BP.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. nsmembrane prote	INC. 43.96% 25.27% 13.10%	CDNA; 3934 BP. Ypeptide PRO1431	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. Ypeptide PRO1431	INC. 43.96% 25.27%
Best Local Similarity: Query Match: RESULT 1299	ID ACD67752 standard; DE CDNA encoding human PN US2003068724-A1.	NTECH 'Y: 'Tity:	0 standa ecreted 68727-A	PD 10-AFK-2003. GETH ) GENENTECH : Percent Similarity: Best Local Similarity: Query Match:	RESULT 1301 ID ACF29244 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN 'US2003068772-A1.	PD 10-APR-2003. PA (GETH ) GENEWIECH Percent Similarity: Best Local Similarity:	ACCOUNT 13702 ID ACD85022 standard; DE Human secreted/trai PN US2003068714-A1.	PD 10-APR-2003.  PA (GETH ) GENENTECH INC.  Percent Similarity: 43.96% Conserva Best Local Similarity: 25.27% Mismacch Procent Mark Mark 12.07% Tadala	RESULT 1303 ID ACD84101 standard; CDNA; DE Human PRO polynucleotide PN US2003068758-A1.	10-APR-2003. (GETH) GENENTECH rcent Similarity: st Local Similarity: ery Match:	KESULT 1304 ID ACD86092 standard; cDNA; 3934 BP. DE Human secreted/transmembrane protein (PRO) PN US2003068776-A1.	PD 10-ARR-2003. PA (GETH) GRENTECH : Percent Similarity: Best Local Similarity: Query Match:	DE ACTOLT9 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003069407-A1.	PD 10-APR-2003, PA (GETH ) GENENTECH : Percent Similarity: Best Local Similarity: Query Match:	RESOLITION ACTIONS TOWA; 3934 BP.  ID ACTIONAL BECTETED POLYPEPTIME PRO1431-encoding PN US2003104555-A1.	05-JUN-2003. (GETH ) GENENTECH ccent Similarity: st Local Similarity:

12		17 39 12		17 339 12	1175.	17 339 12	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 339 12	SEQ ID NO:349.	17 339 12	SEQ ID NO:349.	17 339 12	#175.	17 339 12
Indels:	de #175.	Conservative: Mismatches: Indels:	.de #175.	Conservative: Mismatches: Indels:	ein (PRO) cDNA #	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	(PRO) cDNA	Conservative: Mismatches: Indels:
13.10%	cDNA; 3934 BP. in PRO polypeptide	13.10\$	cDNA; 3934 BP. in PRO polypeptic	INC. 43.96% 25.27% 13.10%	rd; cDNA; 3934 BP. transmembrane prot:	43.96% 25.27% 13.10%	ard; cDNA; 3934 BP. polypeptide PRO1431. 1.	INC. 43.96% 25.27% 13.10%	ard; cDNA; 3934 BP. polypeptide PRO1431. 1.	INC. 43.96% 25.27% 13.10%	ard; cDNA; 3934 BP. polypeptide PRO1431-1.	INC. 43.96% 25.27% 13.10%	ard, cDNA, 3934 BP. polypeptide PRO1431- 1.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. ınsmembrane protein	INC. 43.96% 25.27% 13.10%
Match:	KESULI 130/ ID ACH12042 standard; c DE CDNA encoding human PN US2003049768-A1. PD 13-MAR-2003.	Simil Simil cal Si atch:	1308 H12349 NA enc 200304 -MAR-2	(GETH ) GENENTECH int Similarity: Local Similarity: Match:	KESULT 1309 ID ACD40741 standard; DE Human secreted/tra	3-FEB-2003. tt Similarity: ocal Similarity: Match:	RESULT 1310 ID ACF18213 standard; DE Human secreted poly PN US2003054481-A1.	PD 20-MAR-2003.  PA (GETH ) GENENTECH  PETCENT Similarity:  Best Local Similarity:  Query Match:	RESULT 1311  ID ACP08660 standard; DE Human secreted pol PN US2003049778-A1.	FD 13-MAK-2003.  PETTH GENTH CHEP Percent Similarity: Best Local Similarity: Query Match:	1 stand ecreted 49782-A	FD 13-MAK-2003.  PERCENT SIMILARITY: Best Local Similarity: Ouery Match:	ID ACF52301 standard; DE Human secreted poly US2003054476-A1.	PD (GETH ) GENENTECH INC. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	1314 CD50170 standard; uman secreted/tra S2003068733-A1.	PD 10-APR-2003. PA (GETH) GENENTECH Percent Similarity: Best Local Similarity: Query Match: RESHIT 1315.
Query	NESOLA TO DE O O O O O O O O O O O O O O O O O O	Percent Best Loc Query Ma	<u> </u>	PA (GE Percent Best Loc Query Ma	RESULT TD DE P	PD 13 Percent Best Lo	RESULT ID P DE H	PD 2 PA ( Percen Best I Query	RESULT ID A DE E	PD 13-MA PA (GETH Percent Sir Best Local Querry Match	PESCULT TID BE	PD PA (PERCEN BEST I	TESOLO IN THE PER COLOR	PD PA (Percer Best I	TESOLI ID DE PN	PD 10 PA (G) Percent Best Lo Query M

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PN US20030 PD 13-MAR- PA (GETH )	Percent Similary Result 1324 RESULT 1324	DE BUILD IS SOUSCE PO 13-MONTO PO 13-MONTO PO PO 14-MONTO PO 14-MO	Best Local Query Match: RESULT 1325 ID ACD0393	DE HUMBAN E PN USZ003( PD 27-FEB- Percent Simi	Best Local 5 Query Match RESULT 1326 ID ACD1047	PN USZUGG PD 20-FEB- Percent Simi		PD 27-FEB- Percent Simi Best Local S Query Match:	RESULT 1328 1D ACF4245 1D B Human PN US20036 PD 20-MAR	PA (GETH ) Percent Simi Best Local S Query Match	70.5	PA (GETH ) Percent Simi Best Local & Query Match: PEGHT 1330		PA (GEIH ) Percent Simi Best Local E Ouery Match:	RESULT 1331 ID ACF2181 DE Human E P Human E	PD 13-MAK- PA (GETH ) Percent Simi
SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 39 12	#175.	17 39 12	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	17 339 12	
11-encoding cDNA,	Conservative: Mismatches: Indels:	11-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	11-encoding cDNA,	Conservative: Mismatches: Indels:	(PRO) CDNA	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	11-encoding cDNA,	Conservative: Mismatches: Indels:	ide #175.
cDNA; 3934 BP. lypeptide PRO143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PRO143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PRO143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PRO143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PRO143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. ansmembrane prot	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PR0143	INC. 43.96% 25.27% 13.10%	; cDNA; 3934 BP. lypeptide PRO143	43.96% 25.27% 13.10%	; cDNA; 3934 BP
	PD 10-APK-2003. PD 10-APK-2003. PP (GETH) (BENENTECH Percent Similarity: Best Local Similarity: Cuery Match:	RESOLI 1318 ID ACC26788 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068709-A1.	(GETH) GENENTECH (GETH) GENENTECH scent Similarity: st Local Similarity: ery Match:	RESULT 1317 ID ACF24888 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068716-A1.	PD 10-APR-2003. PA (GETH ) GENENTECH INC. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query_Match: 13.10%	KESULI 1318 ID ACF46468 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding DN 1792013168740.a1	10-APR-2003. (GETH ) GENENTECH coent Similarity: st Local Similarity: sry Match:	RESULT 1319 ID ACP28016 standard; CDNA; 3934 BP. DB Human secreted polypeptide PRO1431-encoding PN VS2003068751-Al.	-2003. ) GENENTECH ilarity: Similarity:	Į,	PD 10.4PR-2003. PA (GETH) GENENTECH Percent Similarity: Best Local Similarity: Query Match:	2 standard ecreted po 73179-A1.	PA (GETH) GENENTECH INC. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10%	RESULT 1322 ID ACF60532 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN U22003087374-A1.	15 to 15	RESULT 1323 ID ACH12656 standard; CDNA; 3934 BP. DE CDNA encoding human PRO polypeptide #175

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499 standard; cDNA; 3934 BP.
secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
3054480-A1.
 secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.3059885-A1.
 310 standard; cDNA; 3934 BP.
secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
3049740-A1.
 818 standard; cDNA; 3934 BP.
secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
3049770-A1.
 914 standard; cDNA; 3934 BP.
secreted/transmembrane protein (PRO) cDNA #175.
3040055-A1.
 472 standard; cDNA; 3934 BP.
secreted/transmembrane protein (PRO) cDNA #175.
3036164-Al.
 114 standard; cDNA; 3934 BP.
secreted/transmembrane protein (PRO) cDNA #175.
3040074-A1.
 079 standard; cDNA; 3934 BP.
secreted/transmembrane protein (PRO) cDNA #175.
3049777-A1.
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 520 standard; cDNA; 3934 BP
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Similarity: 43.968
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39	SEQ	17 39 12	SEO	17 39 12	SEQ.	17 39 12	#175.	17 39 12	#175.	117	, SEQ	17 39 12	#175.	17 39 12	, SEQ	17 39 12
Mismatches: Indels:	4 BP. RO1431-encoding cDNA,	Conservative: Mismatches: Indels:	BP. O1431-encoding cDNA,	Conservative: Mismatches: Indels:	3934 BP. e PRO1431-encoding cDNA.	Conservative: Mismatches: Indels:	BP. protein (PRO) cDNA	Conservative: Mismatches: Indels:	BP. protein (PRO) cDNA	Conservative: Mismatches: Indels:	BP. O1431-encoding cDNA,	Conservative: Mismatches: Indels:	BP. protein (PRO) cDNA	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:
25.27% 13.10%	l; cDNA; 3934 olypeptide PR	43.96% 25.27% 13.10%	cDNA; 3934 BI ypeptide PRO14	INC. 43.96% 25.27% 13.10%	rd; cDNA; 3934 polypeptide PR	INC. 43.96% 25.27% 13.10%		INC. 43.96% 25.27% 13.10%		INC. 43.96% 25.27% 13.10%	cDNA; 3934 ypeptide PR	INC. 43.96% 25.27% 13.10%	cDNA; 3934 nsmembrane ]	INC. 43.96% 25.27% 13.10%	cDNA; 3934 ypeptide PF	INC. 43.96% 25.27% 13.10%
Best Local Similarity: Query Match:	02 standard secreted po 073169-A1.	FD 17-AFK-2003. Percent Similarity: Best Local Similarity: Query Match:	ACF33954 standard; cDNA; DE Human secreted polypeptic PN US2003064457-A1.	PD 03-APR-2003. PA (GETH ) GENENTECH 1 Percent Similarity: Best Local Similarity: Query Match:	standa creted 18711-A1	PD 10-APR-2003. PA (GETH) GENEWTECH 1 Percent Similarity: Best Local Similarity: Query Match:	RESULT 1335  DE Human secreted/transmembrane PN US2003049745-A1.	PA 13-max-2003. PA (GETH ) Percent Similarity: Best Local Similarity: Query Match:	KESULI 1336  ID ACD91161 standard; cDNA; 3934  DE Human secreted/transmembrane by 162001000000000000000000000000000000000	FN 0520009751-A1. PD 13-MAR-2003. PA (GETH ) GENENTECH 1Percent Similarity: Best Local Similarity: GOETY Match:		PD 10-APR-2003. PA (GETH) GENENTECH 1 Percent Similarity: Best Local Similarity: Guery Match:	AESOLI 1393 ID AC087171 standard; cDNA; 393* DE Human secreted/transmembrane PN US2003068777-A1.	(GETH ) GENENTECH rcent Similarity: st Local Similarity: ery Match:	KESOLI 1333 ID ACF60225 standard; CDNA; 3 DE Human secreted polypeptide PN US2003073185-A1. PD 17-APP-2003	ENTECH ty: arity:

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DE Human secret

NGS0056458-A1.

DE Human secret

NGS0056458-A1.

PD 20-MAR-2003.

Percent Similarity: 25.27% Mismatched.

Query Match: 13.10% Indels: 12.

DE Human secreted polypeptide PR01431-encoding cDNA, SEQ ID NO:349.

PN US2003059886-A1.

PD 27-MAR-2003.

PA (GETH) GENERITECH INC.

Conservative: 17

mismatches: 39

Indels: 12

Indels: 12
 ACF08046 standard; cDNA; 3934 BP.

Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003049758-A1.
13-MAR-2003.
NF GENEW GENEWECH INC.
 ACF53836 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003064456-Al.
03-APR-2003.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003096353-A1.
RESULT 1340

ID ACF46775 standard; CDNA; 3934 BP.

DE Human secreted polypeptide PRO1431-encoding CDNA, SEQ ID NO:349.

PN US200308733-A1.

PD 08-MAY-2003.

Percent Similarity: 43.96* Mismatches: 17

Best Local Similarity: 25.27* Mismatches: 39
 ID NO:349
 RESULT 1347
ID ACF40657 standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003064448-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ US2003049772-A1.
 ADA79893 standard; cDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
US2003073173-A1.
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Mismatches:
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Mismatches:
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Indels:
 Indels:
 Query Match: 13.10%
RESULT 1346
ID ACF08353 standard; CDNA; 3934 BP.
 ACF75632 standard; cDNA; 3934 BP.
 PD 13-MAR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 43.96*
Best Local Similarity: 13.10*
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 03-APR-2003.
(GETH) GENENTECH INC.
 DE Human secreted/trans
PN US2003073173-A1.
PD 17-APR-2003
Percent Similarity:
Best Local Similarity: 2
 PA (GETH) GENENTECH 1
Percent Similarity:
Best Local Similarity:
 PA (GETH) GENENTECH :
Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
Query Match:
RESULT 1342
 22-MAY-2003
 Query Match:
RESULT 1343
 Query Match:
RESULT 1348
 Query Match:
RESULT 1341
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Indels:

13.10%

Query Match: RESULT 1357 ID ACF76553 stan DE Human secrete DN 1102003104547	ပိုင်	Best Local Similar Query Match: RESULT 1358 ID ACF61453 stan DE Human secrete	PN US2003096359- PD 22-MAY-2003. PA (GETH) GENEN Percent Similarity	Best Local Similar Query Match: RESULT 1359 ID ACF61760 stan DE Human secrete	PN US2003100061- PD 29-MAY-2003. PA (GETH ) GENEN Percent Similarity	Query Match: RESULT 1360 ID ACD30791 stan DE Human secrete	PN USZU03032125- PD 13-PEB-2003. Percent Similarity		PD 20-MAR-2003. Percent Similarity Best Local Similar	Z Z	PA (GETH ) GENEW Percent Similarity Best Local Similar Query Match:	ID ACF17599 stand BE Human secrete PN US200305460-	PD 20-MAK-2003. PA (GETH ) GENEN Percent Similarity Best Local Similar	Query Match: RESULT 1364 ID ACF07432 stan	PN 03-003-049/553-7 PD 13-MAR-2003. PA (GETH ) GENEN' Percent Similarity	Best Local Similar. Query Match: RESULT 1365 ID ACF20590 stam
17 339 12	#175.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	13 12 12	#175.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	17 39 12	, SEQ ID NO:349.	17 39
Conservative: Mismatches: Indels:	cDNA	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	934 BP. : PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	cDNA	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	3934 BP. de PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	3934 BP. le PRO1431-encoding cDNA,	Conservative: Mismatches:
INC. 43.96% 25.27% 13.10%	cDNA;	INC. 43.96% 25.27% 13.10%	cDNA; 3 ypeptide	INC. 43.96% 25.27% 13.10%	cDNA; 3 ypeptide	INC. 43.96% 25.27% 13.10%	cDNA; 3 ypeptide	INC. 43.96% 25.27% 13.10%	cDNA; 3 nsmembra	INC. 43.96% 25.27% 13.10%	cDNA; 3 ypeptide	43.96% 25.27% 13.10%	cDNA; 3 ypeptide	43.96% 25.27% 13.10%	cDNA; 3 ypeptide	INC. 43.96% 25.27%
PA (GETH ) GENENTECH Percent Similarity: Best Local Similarity: Query Match:	ID ACD47100 standard; cDNA; 3934 BP. DE Human secreted/transmembrane protein (PRO) PN US2003068693-A1.	it de	3 standard ecreted po 68735-A1.	الكهين	RESULT 1351  DE ACHT 1389 standard; cDNA; 3934 BP.  DE Human secreted polypeptide PRO1431-encoding PR US203068753-A1.	5 7 K	ID ACF46161 standard; CDNA; 3934 BP.  E Human secreted polypeptide PRO1431-encoding DN reconsiderals.	PD 10-ARR-2003. PA (GETH ) GENENTECH   Percent Similarity: Best Local Similarity: Query Match:	KENDLI 1333 ID ACD86250 standard; CDNA; 3934 BP. DB Human secreted/transmembrane protein (PRO) PN 192003068765-a1	PD 10-ARR-2003. PA (GETH ) GENENTECH : Percent Similarity: Best Local Similarity: Query Match:	KESULI 135268 standard; CDNA; 3934 BP. DB. Human secreted polypeptide PRO1431-encoding PN US2003082715-A1.	5 7 K	ACCOUNT 1939  ID ACFS-915 standard; cDNA; 3934 BP.  DE Human secreted polypeptide PRO1431-encoding  PN US2003082716-A1.	PD 01-MAY-2003. Percent Similarity: Best Local Similarity: Query Match: RESHIT 13-6	D ACE64908 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068737-A1.	PD 10-APR-2003, PA (GETH ) GENENTECH Percent Similarity: Best Local Similarity:

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Match:	.10%	Indels:	12
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eted	polypeptide PRO1431	-encoding cDNA,	SEQ ID NO:349.
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ity:	.96%	Conservative:	17
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RESULT 1360			
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ilarity:	25.27\$	hes:	39
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13-MAR-2003.			
5	*96	Conservative.	7.1
 cal Similarity:	25.27%	Mismatches:	39
 atch:	10%	Indels:	12
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Percent Similarity: Best Local Similarity: Query Match: RESULT 1374 ID ADA84620 standard DE Human secreted/tr	PN US2003092121-41. PD 15-MAY-2003. PA (GETH ) GENENTECH Percent Similarity:	Best Local Similarity: Query Match: RESULT 1375 ID ACD22264 standard DE Human secreted/tr	PN US2003027276-A1. PD 06-FEB-2003. Percent Similarity:	Best Local Similarity: Query Match: RESULT 1376 ID ACD24611 standard DR Human secreted/fr	PN US2003044920-A1. PD 06-MAR-2003. Percent Similarity:	Best Local Similarity	2	PN US2003027265-A1. PD 06-FEB-2003. Percent Similarity: Best Local Similarity	Query Match: RESULT 1378	1D ACD40121 Standard DE CDNA encoding hum DN HS20030544461-A1	PD 20-MAR-2003. PA (GETH ) GENENTECH	Percent Similarity: Best Local Similarity Onery Match:	RECULT 1379 RECULT 1379 ID ACF13429 standard	DE Human secreted po PN US2003064446-A1.	ğ,	Dest Local Sit Query Match:	2	DE Aumai Secreteu D PN US2003049744-Al. PN 13-MAD-2003	ž č	Query Match: RESULT 1381		
.ve: 17 1: 39	SEQ ID NO:349.	17 339 12	SEQ ID NO:349.	17 339 12	#175.	ţ	39 12	CDNA, SEQ ID NO:349.	:	17 39 10	4	SEQ ID NO:349.	17	39 12	#175.		7.7	39 12			1. 12	
Conservative: Mismatches: Indels:	14 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	3934 BP. de PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	CDNA		Conservative: Mismatches: Indels:	34 BP. PRO1431-encoding cDNA,		Conservative: Mismatches: Indele:		PRO1431-encoding cDNA,	Conservative:	Mismatches: Indels:	CDNA		Conservative:	Mismarches: Indels:	34 BP. Ypeptide #175.		Conservative: Mismatches: Indels:	ACF44609 standard; cDNA; 3934 BP.
INC. 43.96% 25.27% 13.10%	cDNA; 3934 peptide PRC	43.96% 25.27% 13.10%	cDNA; 39 rpeptide	43.96% 25.27% 13.10%	CDNA; 3934 BP semembrane pro	INC.	25.278 13.108	cDNA; 39 Peptide	INC.	43.96* 25.27*	CDNA; 3934 BP	peptide	INC. 43.96%	25.27% 13.10%	cDNA; 3934 nsmembrane p	Ç	43.96%	13.10%	cDNA; 3934 BP n PRO polypept	INC.	43.30% 25.27% 13.10%	CDNA; 39
7 ''	RESULT 1366 ID ACF20897 standard; DE Human secreted poly PN US2003073172-A1.	17-APR-2003. ccent Similarity: it Local Similarity: iry Match:	<pre>f standard; screted poly 73172-A1.</pre>	17-APR-2003. cent Similarity: it Local Similarity: sry Match:	DE AC04714 standard; cDNA; 3934 BP. DE Human secreted/transmembrane protein (PRO) DN 182003068700-21		arıcy: milarity:	RESULT 1369  ID ACR47696 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN 1192003168736-A1.	PD 10-APR-2003. PA (GETH ) GENENTECH I	Percent Similarity: Best Local Similarity: Chorry Match:	RESULT 1370 ID ACF53529 standard;	DE Human secreted polypeptide PRO1431-encoding PN US2003308679-A1.	10-AFR-2003. (GETH ) GENENTECH : ccent Similarity:	τ <b>λ</b> :	RESULT 1371 ID ACD86864 standard; DE Human secreted/trar	PN US2003068767-A1. PD 10-APR-2003.		Best Local Similarity: Query Match: Bestra 1373	KESULI 13/2 ID ACHOSI12 standard; CDNA; 3934 BP. Es CDNA encoding human PRO polypeptide PN US2003073182-Al.	PD 17-APR-2003. PA (GETH ) GENENTECH INC.	Percent Similarity: Best Local Similarity: Query Match:	ID ACF44609 standard;

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rd; cDNA; 3934 BP.
polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
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polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
 rd; cDNA; 3934 BP.
polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
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 RESULT 1393
 ACF11423 standard, cDNA, 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003073171-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003032121-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003064458-A1.
 RESULT 1387
ID ACEZTORO standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003068702-A1.
 ACF24581 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068734-Al.
 #175
 Human secreted/transmembrane protein (PRO) cDNA #175
US2003064464-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175
US2003068719-A1.
 ACD90241 standard, cDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
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 ACD48328 standard; cDNA; 3934 BP
 CDNA; 3934 BP
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 ACD85636 standard; cDNA; 3934 BP
 PN US2003064460-A1.
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Percent Similarity: 43.964
Best Local Similarity: 25.274
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(GETH) GENENTECH INC.
 ID ACF11423 standard; of Human secreted polyy PN US2003073171-A1. PD 17-APR-2003. Percent Similarity: 4 Sest Local Similarity: 4 Query Match:
 DE Human secreted polyr
PN US2003031212-A1.
PD 13-FEB-2003.
Percent Similarity:
 ID ACF24581 standard; of Human secreted polyp.
PN US2003068734-A1.
PD 10-APR-2003.
PA (GETH) GENENTECH IN Percent Similarity:
 ID ACD85636 standard; of Human secreted/trans
PN US2003068719-A1.
PD 10-AFR-2003.
PA (GETH) GENEWIECH IN
Percent Similarity:
Best Local Similarity:
 PN US2003064458-A1.
PD 03-APR-2003.
PA (GBTH) GENENTECH II
Percent Similarity:
Best Local Similarity:
Query Match:
RESULT 1385
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ACF34261 standard;
 ACF50766 standard;
 ACD46486 standard;
 PD 03-APR-2003.
PA (GETH) GENENTECH
Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
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 Query Match:
RESULT 1389
RESULT 1382
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Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003104540-A1.
 ID ACH07316 standard; cDNA; 3934 BP.

DE Human secreted/transmembrane protein (PRO) cDNA #175.

PN US200304972-A1.

PD 13-MAR-2003.

PA (GETH) GENETECH INC.

Percent Similarity: 43.96* Mismacches: 39

Mismacches: 39
 Human secreted/transmembrane protein (PRO) cDNA #175.
US2003049747-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175
US2003049750-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175.
US2003049779-A1.
13-MAR-2003.
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 ACH11428 standard, cDNA, 3934 BP.
CDNA encoding human PRO polypeptide #175.
US2003049766-A1.
 Indels:
 cDNA encoding human PRO polypeptide #175.
US2003049767-A1.
 BP.
 Query Match:
RESULT 1395
ID ACH08237 standard; CDNA; 3934 BP.
 CDNA: 3934 BP
 ACH11735 standard; cDNA; 3934 BP
 RESULT 1398
ID ACH10386 standard; cDNA; 3934 BP
 Query Match: 13.10%
RESULT 1392
ID ACF49231 standard; cDNA; 3934
 ACD83794 standard; cDNA; 3934
Human PRO polynucleotide #175.
US2003068738-A1.
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Local Similarity: 25.27%

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PD 10-APR - BND...
PA (GERH) GEND...
Percent Similarity:
 Query Match:
RESULT 1394
ID ACH07623 standard;
 PN US2003104540-Al.
PD 05-JUN-2003.
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117 122	SEQ ID	17 39 12	ÖES	17 39 12	#175.	17 39 12	#175.	17 39 12	SEQ	17 39 12	SEQ	17 39 12	SEQ.	17 39 12	SEQ.	17 39 12
Conservative: Mismatches: Indels:	encoding cDNA,	Conservative: Mismatches: Indels:	34 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	(PRO) CDNA	Conservative: Mismatches: Indels:	(PRO) CDNA	Conservative: Mismatches: Indels:	934 BP. PRO1431-encoding cDNA,	Conservative: Mismatches: Indels:	1-encoding cDNA,	Conservative: Mismatches: Indels:	34 BP. PRO1431-encoding cDNA	Conservative: Mismatches: Indels:	1-encoding cDNA	Conservative: Mismatches: Indels:
* * * * 0 4 5 0	rd; cDNA; 3934 BP. polypeptide PRO1431	43.96% 25.27% 13.10%	ard; cDNA; 3934 BP. polypeptide PRO1431 1.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. nsmembrane protein	43.96% 25.27% 13.10%	cDNA; 3934 BP. nsmembrane protein	43.96% 25.27% 13.10%	3 Je	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. Ypeptide PRO1431-	INC. 43.96% 25.27% 13.10%	andard; cDNA; 3934 BP. ted polypeptide PRO143 9-Al.	INC. 43.96% 25.27% 13.10%	cDNA; 3934 BP. .ypeptide PRO1431	43.96% 25.27% 13.10%
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Destruction Similarity: 23.278 | Indels: 12 | Destruction State St
 Query Match:
RESULT 1414
ID ACF43995 standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
 DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.

PN US20031044541-A1.

PD 05-JUN-2003

Percent Similarity: 43.96% Conservative: 17

Best Local Similarity: 25.27% Mismatches: 39
 ACP25195 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068712-A1.
 ACF27095 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068730-A1.
 ID NO:349.
ACF38259 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068696-A1.
 ACF76246 standard; cDNA: 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ
10203104545-A1.
05-JUN-2003.
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 ACH06340 standard; cDNA; 3934 BP.
cDNA encoding human PRO polypeptide #175.
US2003049762-A1.
 PN 0.2003.
PD 10-APR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 43.96%
Best Local Similarity: 13.10%
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Percent Similarity: 43.9(
Best Local Similarity: 25.27
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RESULT 1415
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DE HUMAN SEC.
PN US2003068696-AL.
PD 10-APR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 43.9'
Rest Local Similarity: 25.7'
Match: 13.
 PD 10-APR-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 43.

Best Local Similarity: 25.
 US2003104554-A1.
 Query Match:
RESULT 1413
 Query Match:
RESULT 1409
 Query Match:
RESULT 1412
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Query Match: 13.10% Indels: 12
RESULT 1419
ACC93317 standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003032136-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003032133-A1.
 Query Macch: 33
Query Macch: 13.10* Indels: 12
RESULT 1423
ID ACC945Es standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003054467-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003036129-A1.
 RESULT 1422
ID ACF06511 standard; cDNA; 3934 BP.
DB Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003040057-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175
US2003049752-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175 US2003040053-A1.
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 ACHO6647 standard; cDNA; 3934 BP.
cDNA encoding human PRO polypeptide #175.
US2003049765-A1.
 Indels:
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 Indels:
 BB
 ACF19362 standard; cDNA; 3934 BP.
 ACD13053 standard; cDNA; 3934 BP.
 CDNA; 3934
 ACC92703 standard; cDNA; 3934
 DE CDNA encoding human PRO polyr DE CDNA encoding human PRO polyr DN US2003049765-Al.
DD 13-MAR-2003.
PA (GETH) GENENTECH INC.
PACCENT Similarity: 43.96%
Best Local Similarity: 25.27%
PD 13-MAR-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 43.96%

Best Local Similarity: 25.27%
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 PD 13-MAR-2003.
PA (GETH) GENENTECH INC.
Percent Similarity: 43.96
Best Local Similarity: 25.27
Query Match: 13.10
RESULT 1418
 PD 20-MAR-2003.

PA (GRTH) GENENTECH INC.
Percent Similarity: 43.96
Best Local Similarity: 25.27
Query Match: 13.10
RESULT 1424
 RESULT 1417

ID ADA83418 standard; c
DE Human secreted/trans
PN US20049752-A1.
PD 13-MAR-2003.
PA (GETH) GENENTECH IN
 ID ACC92703 standard; of Human secreted polyr. PW US203032133.Al.
PD 13-FEB-2003.
Percent Similarity:
 DE Human secreted polyr
N US2003036129-A1.
PD 20-Pe2-2003.
Percent Similarity: 4
Best Local Similarity: 2
Query Match:
 DE Human secreted/trans
PN US2003040053-A1.
PD 27-EBB-2003.
Percent Similarity:
 DE Human secreted polyr
PN US2003040057-A1.
PD 27-FEB-2003.
Percent Similarity:
 Best Local Similarity:
 DE Human secreted PN US2003032136-AL PD 13-FEB-2003. Percent Similarity:
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ACC97973 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003044932-A1.
06-MAR-2003.
 ACC94238 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003027270-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349. US2003054469-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003059879-A1.
 ACF31768 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003064469-A1.
03-APR-2003.
(GETH) GENENTECH INC.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003049738-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175 US2003032126-A1.
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 cDNA encoding human PRO polypeptide #175.
US2003054463-A1.
 Indels:
 ACD43434 standard; cDNA; 3934 BP.
cDNA encoding human PRO polypeptide #175.
US2003054466-A1.
 PD 13-FEB-2003.
Percent Similarity: 43.96*
Best Local Similarity: 25.27*
ROSULY 1428.
ID ACD43127 standard; CDNA; 3934 BP.
 ACF42192 standard; cDNA; 3934 BP
 ACD31098 standard; cDNA; 3934 BP
 CDNA; 3934 BP
 ACF01696 standard; cDNA; 3934 BP
 ID ACP14964 standard; curr, color DE Human secreted polypeptide PR US2003059879-A1.

PD 27-MAR-2003.

PA (GETH) GRNENTECH INC.

Percent Similarity: 43.96*

Best Local Similarity: 25.27*
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 (GETH) GENENTECH INC.
 ID ACD43127 standard, co
DE CDNA encoding human
PN US2003054465-A1.
PD 20-MAR-2003.
PA (GETH) GENENTECH IN
Percent Similarity: 4
Best Local Similarity: 2
 Query March.
RESULT 1432
TD ACF31768 standard, CF
 Best Local Similarity:
Query Match:
RESULT 1425
 Best Local Similarity:
Query Match:
RESULT 1426
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 PD 06-FEB-2003.
Percent Similarity:
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 20-MAR-2003
 13-MAR-2003
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Query Match:
RESULT 1430
 Query Match:
RESULT 1427
 Query Match:
RESULT 1429
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milarity:

Conservative: 17  Mismatches: 39  Indels: 12  EESULT 1441	ID DE DNG	PD 13-MAR-20	CDNA #175.	PD PA	cDNA #175.	ive: 17 Perce Perce Perce Sest. 39 Query Query 12 RESUL	CDNA, SEQ ID NO:349.	ity st	CDNA, SEQ ID NO:349.	Conservative: 17  Conservative: 17  Mismatches: 39  Indels: 12  Indels: 12  Consty Match: 12  Consty M	CDNA, SEQ ID NO:349.	ve: 17 1: 39	DE Human se PN US200300 PN US200300 PD I3-MAR-2	Conservative: 17  Mismatches: 39  Indels: 12  Conservative: 17  Conservative: 18  Conservative: 18  Conservative: 18  Conservative: 19  Co	CDNA, SEQ ID NO:349.
Percent Similarity: 43.96% Conse Best Local Similarity: 25.27% Misma Questy March: 13.10% Indel	ALSOLI 1433 ID ACD67445 standard; cDNA; 3934 BP. DE CDNA encoding human PRO polypeptide #175 PN US2003064453-A1.	03-APR-2003. (GETH) OBENENTECH INC. (GETH) OBENENTECH INC. st Local Similarity: 25.27% Pry March: 13.10%	ID ACCESS Standard; CDNA; 3934 BP. DE Human secreted/transmembrane protein (PRO) PN US2003064466-a1	03-APR-2003. (GETH ) GENENTECH INC. (GETH ) GENENTECH INC. 43.96* TE LOCAL Similarity: 25.27* 13.10*	About 1993 DE Human secreted/transmembrane protein (PRO) PN US2003064468-A1.	INC. 43.96% 25.27% 13.10%	ID ACF51380 standard; cDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068760-A1.	아 아 아	DD ACF143, Standard, CDNA, 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068769-A1.	INC. 43.96% 25.27% 13.10%	ACF25867 standard; cDNA; 3934 BP. Human secreted polypeptide PRO1431-encoding US2003045700-A1.	PD 06-MAR-2003. PA (GETH) GENENTECH INC. Percent Similarity: 43.96% Conserv Best Local Similarity: 25.27% Mismatc Cuery Match: 13.10% Indels:	KESULY 1439 ID ACF3180 standard; CDNA; 3934 BP. DE Human secreted polypeptide PRO1431-encoding PN US2003068698-A1.	PD 10-APR-2003.  PA (GETH) GENENTECH INC.  EVERTH JAILBAILLY: 43.96% CONSET.  Best Local Similarity: 25.27% Mismat.  OUGTY MAICH: 13.10% Indels	VOLI 1340 F128937 standard; cDNA; 3934 BP. Human secreted polypeptide PRO1431-encoding US2003068759-Al.

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standard, cDNA, 3934 BP.
creted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
8688-A1.
 standard; cDNA; 3934 BP.
reted polypeptide PRO1431-encoding cDNA; SEQ ID NO:349.
552-A1.
 standard; cDNA; 3934 BP.
creted/transmembrane protein (PRO) cDNA #175.
9748-A1.
 standard; cDNA; 3934 BP.
creted/transmembrane protein (PRO) cDNA #175.
8765-A1.
 reted/transmembrane protein (PRO) cDNA #175
 standard; cDNA; 3934 BP.
reted/transmembrane protein (PRO) cDNA #175.
774-A1.
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 standard; cDNA; 3934 BP.
reted/transmembrane protein (PRO) cDNA #175.775-A1.
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oding human PRO polypeptide #175.
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Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003044924-Al.
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 06-MAR-2003
 Query Match:
 Query Match:
RESULT 1464
 Query Match:
RESULT 1462
 Query Match:
 Query Match:
RESULT 1459
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RESULT 1460
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RESULT 1458
 RESULT 1465
 ACF23967 standard, cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068763-A1.
 ALENYOBB Standard; CDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068720-A1.
 Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003068739-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175.
US2003049776-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175 US2003049780-A1.
 Human secreted/transmembrane protein (PRO) cDNA #175
US2003036126-A1.
 ADA78713 standard; CDNA; 3934 BP.
ADA78713 standard; CDNA; 3934 BP.
US2003073181-A1.
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 DE Human secreted/transmembrane protein (PRO) cDNA po 10-APR-2003.

PA (GETH) GENENTECH INC.

Best Local Similarity: 25.27% Mismatches: Query Match: 13.10% Indels:
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PD 10-APR-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 43.96%

Best Local Similarity: 25.27%

Query Match: 13.10% Inder

RESULT 1451

ACF51073 standard; cDNA, 3934 BP.

""" secreted polypeptide PR01431-
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""" an secreted polypeptide PR01431-
 Query Match: 13.10%
RESULT 1457
ID ACC96550 standard; cDNA; 3934 BP.
 ACD11500 standard; cDNA; 3934 BP.
 Query Match:
RESULT 1454
ID ACH09772 standard; cDNA; 3934 BP.
 Query Match: 13.10%
RESULT 1455
ID ACH10693 standard; cDNA; 3934 BP.
 ACD88399 standard; cDNA; 3934 BP
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 Percent Similarity: 43.96%
Best Local Similarity: 25.27%
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(GETH) GENENTECH INC.
11. Similarity: 43.96%
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 RESULT 1449
ID ADA78713 standard, cDNA, DE Human secreted/transmemt PN US2003073181-A1.
PD I7-APR-2003.
PAR (GETH) GENENTECH INC.
Percent Similarity: 43.96
Best Local Similarity: 25.22
 13-MAR-2003.
(GETH) GENENTECH INC.
 PD 13-MAR-2003.

PA (GETH) GENENTECH INC.

Percent Similarity: 43.

Best Local Similarity: 25.
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"1967 standard ...
 Best Local Similarity:
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Percent Similarity:
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RESULT 1450
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CDNA, SEQ ID NO:349.
 RESULT 1461
ID ACF16806 standard; cDNA; 3934 BP.
DE Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PN US2003344073-A1.
 ACC98580 standard, cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003044927-A1.
 ACF41885 standard; cDNA; 3934 BP.
Human secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
US2003040072-A1.
27-FEB-2003.
 ACH04566 standard; cDNA; 3934 BP.
Human cDNA encoding secreted/transmembrane protein PRO1431.
US2003044841-A1.
 DE Human secreted/transmembrane protein (PRO) cDNA #175.
PN US2003032124-A1.
PD 13-FEB-2003
Percent Similarity: 43.96*
Dest Local Similarity: 25.27*
Mismatches: 39
 ACD41355 standard; cDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
US2003064467-A1.
 ACD32326 standard; CDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
US2003054475-A1.
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 Human secreted polypeptide PRO1431-encoding US2003049759-A1.
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 ACF07739 standard; cDNA; 3934 BP
 ACD30484 standard; cDNA; 3934 BP
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Percent Similarity: 43.
Best Local Similarity: 25.
 06-MAR-2003.
(GETH) GENENTECH INC.
 PD 03-APR-2003.
PA (GETH) GENENTECH IN Percent Similarity: 9 Best Local Similarity: 2 Query Match: 1
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 (GETH) GENENTECH
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 PD 06-MAR-2003.
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ID ACF43381 standard; CDNA; 3934 DE Human secreted polypeptide PRO PN US2003104551-A1. PD 05-JUN 2003. PA (GETH ) GENENTECH INC. Percent Similarity: 43.96% Query Macch: 31310% PRENTY 1475.	ID ACHO6033 standard; cDNA; 3934 DB CNNA encoding human PRO polype PN US2003049761-A1. PD 13-MAR-2003. PA (GETH ) GENENTECH INC. Percent Similarity: 43.96% Best Local Similarity: 25.27% Dierry Match:	51 standard, secreted/trar 049757-A1. 22003. GENENTECH J 11arity:		ACSULT 14, 3934  ID ACT10809 standard; CDNA; 3934  DE Human secreted polypeptide PRO PRO US2003036119-A1.  PD 20-FEB-2003.  Percent Similarity: 43.96%  Query Mactoh: Almilarity: 13.10%  RESULT 1479	ID ACC93624 standard; cDNA; 3934 DB Human secreted polypeptide PRO PN US2003036120-A1. PD 20-FEB-2003. Percent Similarity: 43.96% Best Local Similarity: 25.27% Query Match: 13.10% RESULT 1480 TD ACC96243 standard: CDNA: 3934	Iypeptid 1NC. 43.96% 25.27% 13.10%	creted/transmembra 4921-A1. 003. artity: 43.96% milarity: 25.27% milarity: 25.27% milarity: 0.00A, 3 creted polypeptide 9739-A1. GENENTECH INC.
CDNA,	NA, 1	Mismatches: 39 Indels: 12 Indels: 12 I-encoding cDNA, SEQ ID NO:349. Conservative: 17 Mismatches: 39	1 NA,	ь .	<b>м</b> -г	. NA	Mismatches: 39 Indels: 12 Indels: 12 Conservative: 17 Mismatches: 39 Indels: 12
Query Match: 13.10% I RESULT 1466  ID ACF31154 standard; CDNA; 3934 BP. DE Human secreted polypeptide PR01431-PN US2003064455-Al. PD 03-APR-2003. PA (GETH ) GENENTECH INC. PA (GETH ) SARIETTY: 43.96% C	13.10% d; cDNA; 3934 BP. olypeptide PRO1431 H INC.	imilarity: 25.27% imilarity: 25.27% 13.10% 6 standard; cDNA; 3934 BP. ecreted polypeptide PRO1431 73170-A1. 1arity: 43.96% imilarity: 25.27%	13.10% d; cDNA; 3934 BP. olypeptide PRO1433 H INC. 43.96%	: 23.2/4 13.10% d; cDNA; 3934 BP. olypeptide PRO143. H INC. 43.96%	: 25.27% 13.10% d; cDNA; 3934 BP. clectide #175. H INC.	. 25.27% . 13.10% d, cDNA, 3934 BP. olypeptide PRO1431 H INC.	Best Local Similarity: 25.27% Mismatched Query Match: 13.10% Indels: RESULT 1473  ID ACR43074 standard; cDNA, 3934 BP. Indels: Buman secreted polypeptide PR01431-encoding PN US2003104550-A1.  PA (GETH) GENEN'ECH INC. PA (GETH) GENEN'ECH INC. Conservat: Best Local Similarity: 25.27% Mismatched Query Match: I3.10% Indels: RESULT 1474

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4 BP. RO1431-encoding cDNA, SEQ ID NO:349.
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RO1431-encoding cDNA, SEQ ID NO:349.
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Best Loca Query Mat RESULT 14		E C		graph of the	ID ACDS DE Hums PN US20 PD 10-P	0 + 4 E	1D ACF	Percent S Best Loca Query Mat RESULT 14	PN US20 PN US20 PD 13-P	2 # H E	PD TOTAL	Best Loca Query Mat RESULT 10 ID ACDI	PN 20-1 PD 20-1 Percent	Query May Query May RESULT 14 ID ACC DE HUME PN US20	PD 27-F Percent 8 Best Loca	RESULT 14
<i>C</i> 0.01	SEQ ID NO:349.	<i>C</i> 01 <i>C</i> 1	SEQ ID NO:349.	<i>L</i>	SEQ ID NO:349.	<i>7</i>	SEQ ID NO:349.	2 9 7	SEQ ID NO:349.	17 39 12	SEQ ID NO:349.	2	75.	7 6 8 7	75.	ţ
Conservative: 1 Mismatches: 3: Indels: 1:	34 BP. PRO1431-encoding cDNA,	Conservative: 1 Mismatches: 3 Indels: 1	3934 BP. le PRO1431-encoding cDNA,	Conservative: 1 Mismatches: 3 Indels: 1	4	Conservative: 1 Mismatches: 3 Indels: 1	3934 BP. le PRO1431-encoding cDNA, 3	Conservative: 1 Mismatches: 3 Indels: 1	3934 BP. de PRO1431-encoding cDNA, a	Conservative: 1 Mismatches: 3 Indels: 1	3934 BP. le PRO1431-encoding cDNA,	Conservative: 1 Mismatches: 3 Indels: 1	in (PRO) cDNA #17	servative: 1 matches: 3	sin (PRO) cDNA #17	
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Percent Similarity: Best Local Similarity: Query Match:	25 standa secreted 059882-A1 -2003	PA (GETH ) GENENTECH Percent Similarity: Best Local Similarity: Query Match:	39 standa secreted 059884-A1	2003. GENENTECH larity: imilarity:	7 standard ecreted po- 68687-A1.	10-APR-2003. (GETH ) GENENTECH : cent Similarity: Local Similarity: Y Match:	ACF33303 standard; Human secreted pol US2003073186-A1.	PD 17-APR-2003.  PA (GETH) GENENTECH I Percent Similarity: Best Local Similarity: Query Match:	resold 148/ ID ACF547 standard; cDNA; 39 DE Human secreted polypeptide PN US200306443-A1.	PD 03-APR-2003. PA (GETH ) GENENTECH IN Percent Similarity: 9 Best Local Similarity: 2 Query Match: 1	ACF48617 standard; cDNA; 33 ACF48617 standard; cDNA; 35 Human secreted polypeptide US2003064444-A1.	PD 03-APR-2003. PA (GETH) GENENTECH 1 Percent Similarity: Best Local Similarity: Cuery Match:	ACD47407 standard; cDNA; 3934 BP. Human gecreted/transmembrane protein	PD 10-APR-2003. PA (GETH ) GENENTECH PErcent Similarity: Query Match:	RESULT 1490 ID ACD49249 standard, CDNA, 3934 BP. En Human secreted/transmembrane protein PN US2003068710-A1.	PD 10-APR-2003. PA (GETH ) GENENTECH

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man secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
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ETH) GENENTECH INC.
Similarity: 43.96*
Conservative: 17
Mismatches: 39
atch: 13.08*
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FOO581 standard; cDNA; 3934 BP.
man secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
 F30165 standard; cDNA; 3934 BP.
nan secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
Q03073178-A1.
APR-2003.
 762067 standard; cDNA; 3934 BP.
nan secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
2003104538-A1.
 1498
299187 standard; cDNA; 3934 BP.
nan secreted polypeptide PRO1431-encoding cDNA, SEQ ID NO:349.
PEB-2003.
 LT 1493
ACD87478 standard; CDNA; 3934 BP.
ACD87478 standard; CDNA; 3934 BP.
Human secreted/transmembrane protein (PRO) cDNA #175.
US2003068774-A1.
10-APR-2003.
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(GETH) GENENTECH INC.
Conservative: 17
t Local Similarity: 25.27% Mismatches: 39
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##11000 standard; CDNA; 3934 BP.
##11000 setected/transmembrane protein (PRO) CDNA #175.
2003049781-A1.
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Percent Similarity: 25.27% Mismatches: 17

Best Local Similarity: 25.27% Mismatches: 39

Query Match: 13.10% Indels: 12

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Scoring table:

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/clone lib="NIH MGC 184"
/clone lib="NIH MGC 184"
/note="Organ: Pooled-Glandular; Vector: pDNR-LIB; Site_1:
Sfil (ggccattatggcc); Site_2: Sfil (ggccgctcggcc);
Library is oligo-dT primed and directionally cloned. CDNR
was prepared from a glandular pool of tissues from thyoid,
parathyroid, adrenal, cortex and pineal gland. 5' and 3'
adaptors were used in cloning as follows: 5' adaptor
sequence: 5'-CACGGCGATTATGGCC-3' and 3' adaptor sequence:
5'-ATTCTAGAGGCGGGGGGGCGATG-GT(30)BN-3' (where B = A,
C, or G and N = A, C, G, or T). Average insert size 1.38
kb (range 0.60-3.5 kb): 15/15 colonies contained inserts
by PCR. This library was enriched for full-length clones
and was constructed by Clontech Laboratories (Palo Alto,
CA). Note: this is a NIH_MGC Library."
 Email: cgapbs-remail.nih.gov
Tissue Procurement: Dr. Michael Brownstein and Dr. Miklos Palkovits
CDNA Library Preparation: CLONTECH Laboratories, Inc.
CDNA Library Parayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Agencourt Bioscience Corporation
Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at:

http://image.llnl.gov
Plate: NDCM151 row: e_column: 19
 CB958894 800 bp mRNA linear EST 29-APR-2003
AGENCOURT 13785021 NIH MGC_184 Homo sapiens cDNA clone
IMAGE:30352338 5', mRNA sequence.
 448
 332
 272
 447 ITCAGNAAGC---ACATTGNAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAG 391
 9
 80
 40
 Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi, Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
11 (bases 1 to 800)
NHH-MGC http://mgc.nci.nih.gov/.
National Institutes of Health, Mammalian Gene Collection (MGC) Unpublished (1999)
 rGlyLysGlyIleValLysGlyArgAsnLeuAspSerArgGlyLeuIleLeuGlyAlaGl
 AGGCAAGGGCATAGTTAAAGGACGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGA
 MetThrPhePheLeuSerLeuLeuLeuLeuValCysGluhlaIleTrp-ArgSerAs
 507 ATGACATTTTTCGGTCACNATTATNATGGTTGGTATGTGAAGCTATTTGGNAGATCCAA
 nSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHisSe
 rGlnProThrGlnSerSerLeuGluAspSerValThrProThrLysAlaValLysThrTh
 271 AGCCTGGGGCAGGGTGTAAAGAAAACACT 241
 80 uAlaTrpGlyArgGlyValLysLysAsnThr 90
 Contact: Robert Strausberg, Ph.D.
 'organism="Homo sapiens"
Gaps:
 High quality sequence stop: 488.
Location/Qualifiers
 US-09-989-293A-377 (1-90) x AU185777 (1-663)
 /mol_type="mRNA"
/db_xref="taxon:9606"
 CB958894.1 GI:30215010
 Homo sapiens (human)
Homo sapiens
 1. .800
 CB958894
 EST.
 20
 40
 390
 9
 331
 VERSION
KEYWORDS
SOURCE
ORGANISM
 LOCUS
DEFINITION
 REFERENCE
AUTHORS
TITLE
JOURNAL
COMMENT
 ACCESSION
 RESULT 2
CB958894
 FEATURES
 ORIGIN
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Eukaryofe, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
1 (bases I to 659)
Dias Neto, E., Garcia Correa, R., Verjovski-Almeida, S., Briones, M.R., Nagai, M.A., da Silva, W. Jr., Zago, M.A., Bordin, S., Costa, F.F., Goldman, G.H., Carvalho, A.F., Matsukuma, A., Baia, G.S., Simpson, D.H., Brunstein, A., deoliveira, P.S., Bucher, P., Jongeneel, C.V., Soares, F., Brentani, R.R., Reis, L.F., G. Souza, S.J. and Simpson, A.J.
 BIO18962
IL3-MT0267-050101-437-A02 MT0267 Homo sapiens cDNA, mRNA sequence.
BIO18962
 Fax: +55-11-2707001
Email: asimpson@ludwig.org.br
This sequence was derived from the FAPESP/LICR Human Cancer Genome
Project. This entry can be seen in the following URL
(http://www.ludwig.org.br/scripts/gethtml2.pl?tl=IL3&t2=IL3-MT0267-050101-437-A02&t3=2001-01-05&t4=1)
 461
 462 chacceachearcarchthagaagacagrafachectactaccaaagcrafcaaaacaac 521
 582 TGTTATTCTTATTCCAGCAATTGTCCACTAAAATTTCCCTGGGGGATTGGGAAGATAAAA 641
 40
 9
 74
 Contact: Simpson A.J.G.
Laboratory of Cancer Genetics
Ludwig Institute for Cancer Research
Rua Prof. Antonio Prudente 109, 4 andar, 01509-010, Sao Paulo-SP,
 Shotgun sequencing of the human transcriptome with ORF expressed
 TCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAGT
 SerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHisSer
 GlnProThrGlnSerSerLeuGluAspSerValThrProThrLysAlaVal-LysThrTh
 rGly------LysGlyIleValLys------GlyArgAsnLeuAspSerArgGl
 Sequence tags
Proc. Natl. Acad. Sci. U.S.A. 97 (7), 3491-3496 (2000)
20202663
 642 AAGAACAATTGGGCTTGGGGCCAAACCTTGGGGGGCCTTCCCTAAAA 687
 87
 800
70
10
29
 Matches:
Conservative:
Mismatches:
 Indels:
 (1-800)
 Seq primer: puc 18 forward
High quality sequence stop: 639.
Location/Qualifiers
 US-09-989-293A-377 (1-90) x CB958894
 BI018962.1 GI:14425592
 1.15e-26
306.00
66.38%
60.34%
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1. .663
Plate: 30 row:
Seq primer: T7
Class: BAC ends.
 Homo sapiens
 Percent Similarity:
Best Local Similarity:
 82 TrpGly 83
 TGGGGT
 Alignment Scores:
 EST.
 4
 12
 Query Match:
 source
 KEYWORDS
SOURCE
ORGANISM
 Pred. No.:
 DEFINITION
 ACCESSION
VERSION
 REFERENCE
AUTHORS
 TITLE
JOURNAL
COMMENT
 RESULT 5
AV721179
 FEATURES
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 663 bp DNA linear GSS 12-MAY-2000 RPCI-23-30A15.TV RPCI-23 Mus musculus genomic clone RPCI-23-30A15, AZ121459
 /db xrefe="taxon:9606"
/dev stage="Adult"
/clone_lib="wyr026"
/clone_lib="wyr026"
/note="Organ: marrow; Vector: puc18; Site_1: SmaI; Site_2: SmaI; A mini-library was made by cloning products derived from ORESTES PCR (U.S. Letters Patent application No. 196,716 - Ludwig Institute for Cancer Research) profiles into the pUC 18 vector: Reverse transcription of tissue mRNA and cNNA amplification were performed under low
 Zhao,S., Nierman,W., Feldblyum,T., Malek,J., Shatsman,S.,
Akinret,B., Levins,M., Mcgann,S., Tsegaye,G., Geer,K., Krol,M., de
Jong,P. and Fraser,C.M.
Mouse BAC End Sequences from Library RPCI-23
Unpublished (1999)
 Email: szhoo@tigr.org
Clones are derived from the mouse BAC library RPCI-23. For BAC
Library availability, please contact Pieter de Jong
(pieter@dejong.med.buffalo.edu. Clones may be purchased from
BACPAC Resources (http://bacpac.med.buffalo.edu/orderingframe.htm)
or from Resea ch Genetics (info@resgen.com). BAC end page.
http://www.tigr.org/tdb/bac_ends/mouse/bac_end_intro.html
 558
 557 AATTCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCAC 498
 497 AGTCAACCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTGAAAACC 438
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 663)
 39
 59
 ---GluAlaIleTrpArgSer
 20 AsnSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHis
 40 SerGlnProThrGlnSerSerLeuGluAspSerValThrProThrLysAlaValLysThr
 Other_GSSS: RPCI-23-30A15.TJ
Contact: Shaying Zhao
Debarment of Bukaryolic Genomics
The Institute for Genomic Research
7712 Medical Center Dr., Rockville, MD 20850, USA
Pal: 301 838 0200
Pax: 301 838 0208
 Conservative:
Mismatches:
Indels:
 Length:
Matches:
 1. .659
/organism="Homo sapiens"
 Gaps:
 LeuSerLeuLeuLeuLeuValCys----
 US-09-989-293A-377 (1-90) x BI018962 (1-659)
 stringency conditions.
 Mus musculus (house mouse)
 mol_type="mRNA"
 AZ121459.1 GI:7788387
 1.2e-19
247.50
87.10%
 80.65%
53.57%
 Mus musculus
 ACAGGG 432
 60 ThrGly 61
 Similarity:
 Percent Similarity:
 Alignment Scores:
 Query Match:
DB:
 source
 RESULT 4
AZ121459/c
 LOCUS
DEFINITION
 Best Local
 . No. .
 ORGANISM
 TITLE
JOURNAL
COMMENT
 ACCESSION
 VERSION
KEYWORDS
SOURCE
 REFERENCE
 AUTHORS
 DRIGIN
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/lab host="DHIOB"
/clone lib="RPCI-23"
/clone lib="RPCI-23"
/note="Corgan: Kidney/Brain; Vector: pBACe3.6; Site_1:
ECORI; Site 2: EcoRI; Female C57BL/6J mouse kidney and/or
brain genomIc DNA was isolated and partially digested
with a combination of EcoRI and EcoRI Methylase. Size
selected DNA was cloned into the pBACe3.6 vector at the
ECORI sites. The ligation products were transformed into
DHIOB electrocompetent cells (BRL Life Technologies). "
 EST 16-OCT-2000
 To (bases 1 to 673)

Gu,Y., Peng,Y., Song,H., Huang,Q., Yang,Y., Gao,G., Xiao,H., Xu,X.,
Li,N., Qian,B., Liu,F., Qu,J., Gao,X., Cheng,Z., Xu,Z., Zeng,L.,
Xu,S., Gu,W., Tu,Y., Jia,J., Fu,G., Ren,S., Zhong,M., Lu,G., Hu,R.,
Chen,J., Chen,Z. and Han,Z.

Homo sapiens cDNA HTB clones
Unpublished (2000)
 248 TTTCTGTCACTATTGCTGTTCGTGACGTATGAAGCATTTTGGTGACAATTCAGGGAGA 189
 43
 63
 72
 81
 13
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
 Contact: Zeguang Han
Chinese National Human Genome Center at Shanghai
351 Guo Shoujing Road, Zhangjiang Hi-Tech Park, Pudong, Shanghai
 AV721179

AV721179 HTB Homo sapiens cDNA clone HTBAKF11 5', mRNA sequence. AV721179

AV721179.1 GI:10818331
 PheleuSerLeuLeuLeuLeuValCysGluAlalleTrpArgSerAsnSerGlySer
 24 AsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHisSerGlnProThr
 71 AGCACGAAAGGAGC-AATGGGATCCCGGATTCCAGAGGTTCATTTCCTGGTGCTGAGGCC
 131 GAATCATCTTTAGATGAGAAGGTGGCTCCCAAGGCATCCCAAACTACAGGCAAGGGC
 64 IleVallySGlyArgAsn----LeuAspSerArgGlyLeuIleLeuGlyAlaGluAla
 44 GlnSerSerLeuGluAspSerValThrProThrLysAlaValLysThrThrGlyLysGly
 663
50
7
22
4
 Conservative:
Mismatches:
Indels:
 Matches:
 Length:
 /organism="Mus musculus"
 /mol_type="genomic DNA"
/strain="C57BL/6J"
 US-09-989-293A-377 (1-90) x AZ121459 (1-663)
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Location/Qualifiers
 /sex="Female"
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209.00
69.51%
60.98%
45.24%
 Homo sapiens (human)
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855 bp mRNA linear EST 26-JUN-2001
602891529F1 NCI_CGAP_Lu29 Mus musculus cDNA clone IMAGE:5036647 5',
mENA sequence.
 Eukaryota, Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus. 1 (bases 1 to 855)
NIH-MGC http://mgc.nci.nih.gov/.
National institutes of Health, Mammalian Gene Collection (MGC)
Unpublished (1999)
 274 CTGTTGAAGAGTGACAGCTTTCCATCAAGAAATAAAGACAACCAAGAGTCAACCCACACAA 333
 Contact: Robert Strausberg, Ph.D.
Email: cgapbs-r@mail.nih.gov
Tissue Procurement: Gilbert Smith, Ph.D.
Tissue Procurement: Gilbert Smith, Ph.D.
CDNA Library Preparation: Life Technologies, Inc.
CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at:
http://image.llnl.gov
Plate: LLAMAIIOI row: j column: 08
High quality sequence start: 27
High quality sequence stop: 825.
Location/Qualifiers
 214 cregrearaacreregrecreagraceregerarrregagarecagricagegaacaac 273
 /clone lib="NCI_CGAP_Lu29"
/note="Organ: lung; Vector: pCWV-SPORT6; Site 1: Sall;
Site_2: Not1; cloned unidirectionally. Primer: Oligo dT.
Library constructed by Life Technologies. Investigator
providing samples: Gilbert Smith, NIH"
 /db_xref="taxon:10090"
/clone="INAGE:5036647"
/tissue_type="spontaneous tumor, metastatic to mammary.
Stem cell origin."
/lab_host="DH108"
/note="Vector: pcDNA3.1; Site 1: EcoRI; Site 2: NotI; Library made with RNA pooled from multiple tissues including liver, lung, hypothalamus, pituitary, and placenta/endometrium."
 5 LeuSerLeuLeuLeuLeuValCysGluAlalleTrpArgSerAsnSerGlySerAsn
 25 ThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHisSerGlnProThrGln
 61
 SerSerLeuGluAspSerValThrProThrLysAlaValLysThrThrGly
 582
35
9
13
0
 Conservative:
Mismatches:
Indels:
 Length:
Matches:
 organism="Mus musculus"
 US-09-989-293A-377 (1-90) x CB420818 (1-582)
 /mol_type="mRNA"
/strain="CZECH II"
 Mus musculus (house mouse)
Mus musculus
 BI107684.1 GI:14558577
 23e-11
 174.00
77.198
61.408
37.668
 1. .855
 Best Local Similarity:
Query Match:
 Percent Similarity:
 Alignment Scores:
 45
 VERSION
KEYWORDS
SOURCE
ORGANISM
 No.:
 DEFINITION
 AUTHORS
TITLE
JOURNAL
COMMENT
 ACCESSION
 REFERENCE
 BI107684
 FEATURES
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 Contact: Smith TPL
USDA, ARS, US Meat Animal Research Center
PO Box 166, Clay Center, NE 68933-0166, USA
Tel: 402 762 4396
Fax: 402 762 4390
Bmail: smithDeemall.marc.usda.gov
Bmail: smithDeemall.marc.usda.gov
trimmed with the aid of the trim_alt option. Vector identified with
cross match v0.990329.
Plate: FQY8014 row: B column: 7
Seg primer: GTANTACGACTCACTATAGGG.
 Eukaryotta; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae; Bovinae; Bos. 1 (bases 1 to 582) Smith, T.P.L., Roberts, A.J., Echternkamp, S.E., Chitko-McKown, C.G., Wray, J.E. and Keele, J.W. A second set of bovine ESTs from pooled-tissue normalized libraries Unpublished (2003)
 EST 25-MAR-2003
 2 GGAAGCAACACATTGGAGAATGGCTACTTCTATCAAGAAATAAAGAGAACCACAGTCAA 61
 /clone lib="HTB"
/note="Vector: pBluescript sk(-); Site_1: EcoRI; Site_2:
XhoI"
 22 GlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLysGluAsnHisSerGln 41
 61
 42 ProThrGlnSerSerLeuGluAspSerValThrProThrLysAlaValLysThrThrGly
 CB420818 582 bp mRNA linear 593806 MARC 6BOV Bos taurus cDNA 5', mRNA sequence.
 673
40
0
0
0
 201203, P. R. China
Tel: 86-21-50801919(ex.45)
Fax: 86-21-50801925.
Email: hanzg@chgc.sh.cn
This clone is available at CHGC in Shanghai.
Location/Qualifiers
 Conservative:
Mismatches:
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/dev_stage="Adult"
/lab_host="SOLR"
 Length:
Matches:
 Indels:
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/mol_type="mRNA"
/db_xref="taxon:9606"
 Gaps:
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/db_xref="taxon:9913"
 /tissue type="pooled"
/lab host="DH10B"
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 Location/Qualifiers
1. .582
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 CB420818
CB420818.1 GI:29187264
 100.00%
 44.59%
 i. .673
 Bos taurus (cow)
 Best Local Similarity:
 Bos taurus
 Percent Similarity:
 Alignment Scores:
 Query Match:
DB:
 DEFINITION
ACCESSION
VERSION
KEYWORDS
SOURCE
ORGANISM
 Pred. No.:
 TITLE
JOURNAL
COMMENT
 RESULT 6
CB420818
 REFERENCE
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CL005921 531 bp DNA linear GSS 29-DEC-2003
ZMMBBb0540E16f ZMMBBb (HindIII) Zea mays genomic clone
ZMMBBb0540E16 5', genomic survey sequence.
 .88 CTGCTGTCGTCCTCTCCCCGGAGCCCAAACACCAGAGGATTCTTGCTCTCTCACTCCAGA 247
 308 ACGCCCACCGTCAGATTGGCAAAGGCCGCAATCGCACCTCCTC-----GAG 355
 The Plant Genome Initiative at Rutgers, Waksman Institute, Rutgers
University
 Eukaryóta, Viridiplantae, Streptophyta, Embryophyta, Tracheophyta, Spermatophyta; Magnoliophyta, Liliopaida, Poales, Poaceae, PACCAD clade, Panicoideae, Andropogoneae, Zea.
 --AspSerValThrProThrLysAlaValLys 58
 ThrThrGlyLysGlyIleValLysGlyArgAsnLeuAspSerArgGlyLeuIleLeuGly 78
 -----LeuLeuLeuValCysGluAlaIle 16
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/cultivar="B73"
/db xref="taxon.4577"
/clone="zMMBBD 540E16"
/lab host="E. col! BH.08"
/clone lib="zMMBBD (HindIII)"
/note="Vector: pCUGI; Site_1: HindIII"
 17 TrpArgSerAsnSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLys
 Bharti, A.K., Young, S., Kavchok, S., Keizer, G., Bronzino, A.C., Zohovetz, V., Fuks, G., Yu, Y., Wing, R. and Messing, J. Sequencing of the maize genome at PGIR (2003c) Curpublished (Unpublished Act.) A.K.
Dr. Joachim Messing's lab
 Onto Tail of Manager Road, Piscataway, NJ 08854, USA Tel: 732 445 3801 Fax: 732 445 5735 Fax: 732 445 575 Fax:
 531
31
9
20
36
5
 Conservative:
Mismatches:
Indels:
 Matches:
 Length:
 High quality sequence start: 56.
Location/Qualifiers
1. .531
/organism="Zea mays"
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 42 ProThrGlnSerSerLeuGlu----
 356 ACAGAGCAGGGGGGCAAAGGT 376
 ThrPhePheLeuSerLeu-----
 79 AlagiuAlaTrpGiyArgGiy 85
 CL005921.1 GI:40367199
 (bases 1 to 531)
 84.00
41.67%
32.29%
18.18%
 Class: BAC ends
 Percent Similarity:
Best Local Similarity:
 Zea mays
 Zea mays
 CL005921
 Alignment Scores:
 59
 ~
 Query Match:
 SOURCE
ORGANISM
 source
 LOCUS
 Pred. No.:
 TITLE
JOURNAL
COMMENT
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KEYWORDS
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 CL005921
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 BJ675325 HCEST library Haplochromis chilotes cDNA clone no152h09,
 Haplochromis chilotes
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
Acanthomorpha; Acanthopterygii; Percomorpha; Perciformes;
Labroidei; Cichlidae; Haplochromis.
1 (bases 1 to 621)
 15 AlaIleTrpArgSerAsnSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArg 34
 Watanabe,M., Kobayashi,N., Shin-i,T., Kohara,Y. and Okada,N. Orf sequences of cichlid in Lake Victoria are essentially same Unpublished (2004)
 ------AsnHisSerGln
 855
28
7
11
1
 621
29
12
30
16
 Center For Genetic Resource Information
National Institute of Genetics
National Anstitute of Genetics
Tel: Wishima, Shizuoka 411-8540, Japan
Tel: 81-559-81-6856
Fax: 81-559-81-6855
Email: tshini@genes.nig.ac.jp.
Location/Qualifiers
 /organism="Haplochromis chilotes"
/mol_type="mRNA"
/db_xref="taxon:257977"
 Length:
Matches:
Conservative:
Mismatches:
Indels:
 Length:
Matches:
Conservative:
Mismatches:
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/tissue_type="jaw"
/dev_stage="varied"
/clone_lib="HCEST library"
 Indels:
 Gaps:
 Gaps:
 US-09-989-293A-377 (1-90) x BJ675325 (1-621)
 US-09-989-293A-377 (1-90) x BI107684 (1-855)
 31 PheLeuSerArgAsnLysGlu-----
 AAGGCATCCCAAACTACAGGA 448
 LysAlaValLysThrThrGly 61
 BJ675325.1 GI:46516089
EST.
 3.23e-06
138.50
74.47%
59.57%
29.98%
 Haplochromis chilotes
 2.29
89.00
47.13%
33.33%
 mRNA sequence.
 Percent Similarity:
Best Local Similarity:
 Percent Similarity:
Best Local Similarity:
 BJ675325
Alignment Scores:
 Alignment Scores:
 55
 128
 Query Match:
 Query Match:
DB:
 LOCUS
 Pred. No.:
 Pred. No.:
 ORGANISM
 ACCESSION
VERSION
KEYWORDS
SOURCE
 AUTHORS
TITLE
JOURNAL
COMMENT
 REFERENCE
 RESULT 8
BJ675325
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/mol_type="mRNR" crp. coll line"

/db_xref="laxon:9666"

/db_xref="laxon:9666"

/lone="lMAGE:4454450"

/tissue_Lype="hypernephroma, cell line"

/lab_host="DHIOB (phage-resistant)"

/lab_host="DHIOB (phage-resistant)"

/olone="lib="NIH MGC B9"

/note="organ: kidney; Vector: pCMV-SPORT6; Site_1: NotI;

Site_2: Sall; Cloned unidirectionally; oligo-dT primed.

Average insert size l.3 kb. Library enriched for full-length clones and constructed by Life Technologies.

Note: this is a NIH_MGC Library."
 BG165693 924 bp mRNA linear EST 06-FEB-2001
602344512F1 NIH_MGC_B9 Homo sapiens cDNA clone IMAGE:4454450 5',
 Unpublished (1999)
Contact: Robert Strausberg, Ph.D.
Email: cgapbs-remail.nih.gov
Tissue Procurement: ATC
CDNA Library Preparation: Life Technologies, Inc.
CDNA Library Preparation: Life Technologies, Inc.
CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at:
http://image.llnl.gov
http://image.llnl.gov
Plate: LLAM10246 row: h column: 03
High quality sequence stop: 703.
High quality sequence stop: 703.
717 AGGAAGATCTTGAAAGGGAAATTTÁAÁACAAGGGCATTTGGAÁTCTAGAÁGGGGTTTAAG 776
 534 riccicadecrerecrerentenearecrecrecaecaecarecrerriderreagaic 475
 Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi, Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo. I (Dases 1 to 24)
NIH-MGC http://mgc.nci.nih.gov/.
 17 TrpArgSerAsnSerGlySerAsnThrLeuGluAsnGlyTyrPheLeuSerArgAsnLys 36
 --CysGluAlaile
 -AlaValLysThrThrGlyLysGlyIleValLysGlyArgAsn
 777 gegeenceaceesakeengarireesesekagaraarreeseseka 821
 84
 LeuAspSerArgGlyLeuIleLeuGlyAlaGluAlaTrpGlyArg
 Conservative:
 Mismatches:
 Length:
Matches:
 Indels:
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 474 CTCACATTCCATCTTCATCAGAATCAATA-
 (1-924)
 4 PheleuSerLeuLeuLeuLeuLeuVal --
 US-09-989-293A-377 (1-90) x BG165693
 BG165693.1 GI:12672396
 Homo sapiens (human)
 27.4
82.00
40.00$
27.27$
17.75$
 .924
 mRNA sequence.
 Homo sapiens
 Best Local Similarity:
 BG16569.
 Percent Similarity:
 Alignment Scores:
 56
 70
 Query Match:
DB:
 source
 ..
ON
 BG165693/c
 DEFINITION
 ORGANISM
 AUTHORS
TITLE
JOURNAL
 ACCESSION
 VERSION
KEYWORDS
 REFERENCE
 FEATURES
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 SOURCE
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 3' adaptor sequence:
5'-ATTCTAGAGGCGGCGGAGATG-GT(30)BN-3' (where B = A,
C, or G and N = A, C, G, or T). Average insert size 1.55
kb (range 0.9-4.0 kb). 15/15 colonies contained inserts
by PCR. This library was enriched for full-length clones
and was constructed by Clontech Laboratories (Palo Alto,
CA)."
 833 bp mRNA linear EST 27-SEP-2000 61671978F1 NIH_MGC_S3 Homo sapiens cDNA clone IMAGE:3960715 5', BE865626
 /tissue type="arcinoma, cell line"
/lab host="DH108 (T1 phage-resistant)"
/clone_lib="NIH MGC 53"
/note="Organ: bladder, Vector: pDNR-LIB (Clontech);
Site_1: Sfil (ggcgctcgggcc); Site_2: Sfil
(ggcattatggcc); Double-stranded cDNA was prepared from cell line RNA. 5' and 3' adaptors were used in cloning as follows: 5' adaptor sequence: 5'-CACGGCCATTATGGCC-3' and
 73
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
I (bases 1 to 833)
NIH-MGC http://mgc.nci.nih.gov/.
 53
 Unpublished (1999)
Contact: Robert Strausberg, Ph.D.
Email: cgapbs.remail.nih.gov
Tissue Procurement: ATCC
CDNA Library Preparation: CLONTECH Laboratories, Inc.
CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MCC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at: image.llnl.gov
Plate: LLCM843 row: K. column: 20
 -----AsnHisSerGlnProThrGlnSerSerLeuGluAspSerValThrPro
 54 ThrLysAlaValLysThrThrGlyLysGlyIleValLysGlyArgAsnLeuAspSerArg
 411 --- Grgarchiaegeeceaegeecieegaaaareeaeireerreeaar 455
 89
 74 GlyLeuIleLeuGlyAlaGluAlaTrpGlyArgGlyValLysLysAsn
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1
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Conservative:
Mismatches:
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82.00
50.91%
43.64%
17.75%
 . .833
 Homo sapiens
 Query Match:
DB:
 Percent Similarity:
 37 Glu-
 Alignment Scores:
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VERSION KEYWORDS SOURCE ORGANISM

TITLE JOURNAL COMMENT REFERENCE AUTHORS

FEATURES

DEFINITION

ACCESSION

RESULT 10

BE865626

51

37 GluAsnHisSerGlnProThrGlnSerSerLeuGluAspSerVal-------

à

36 LysGluAsnHisSerGlnProThrGlnSerSerLeuGluAspSerValThrProThrLys 55

ð

No.:

Score: Pred.

ORIGIN

16

Query Match: 17.53% Indels: DB: 6aps:	US-09-989-293A-377 (1-90) x CNS04H30 (1-1121)	73	Oy 18 ArgSerAsnSerGlySerAsn :::		859 39	Db 919 TCAGTCCACAGTTCTCCTCCAGCTATGACACTT  Oy 54 ThrLysAlaValLysThrThrGlyLysGlyIle  Db 979 ACCAGTTTTTTGGTCTGCGGGAAGTAGGATA	Qy 74 GlyLeuIleLeuGly 78 Db 1039 AACTTTGACCTCGGG 1053	RESULT 13 B1914658/c LOCUS DEDING	MENSA SEQUENCE.  BI914658  BI914658.1 GI:16199003	EST. Homo sapiens (human) Homo sapiens	Duranjous, mecazoa, Chordata, Crana Mammalia: Eutheria: Primates; Catar: REFERENCE 1 (bases 1 to 683)		Email: cgapbs-r@mail.nih.gov Tissue Procurement: Life Technologi CDNA Library Preparation: Life Tec	CONN LIDITAL ALLAYED BY: INC 1.W.A DNA Sequencing by: Incyte Genomics, Clone distribution: MGC clone distribution: MGC clone distribution found through the I.M.A.G.E. Consort		Cource 1683 Organism="Homo.gapiens" (organism="Homo.gapiens"	/ MOCYPB=_MXXM / MDCYPB=_MXXM / Clone="Laxon:9606" / Clone="Laxon:9606" / Alone = "DHLOB" / Alone = This MYC	// / / / / / / / / / / / / / / / / / /	and male age 26 weeks. Lift directionally cloned (EcoR) cloning). Average insert so 0.7-3.5 kb. Library is not full length clones and was
	ThrProThrLysAlaValLysThrThrGlyLysGlyIleValLysGlyArgAsn 69 	eLeuGlyAlaGluAla	ICTCACTCGGGTCTCGTCGTGGGTCCCACGGCCACCAGTACCCACTGCCACGACATT 280 		CNSO4H30 Tetraodon nigroviridis genome survey sequence T7 end of clone	JONIS OI LIBRARY G FROM TETRACOGON NIGROVIRIDIS, GENOMIC SURVEY SEQUENCE. AL290421 AL290421.1 GI:8029001 GSS; genome survey sequence. Tetracodon nigroviridis	Tetraodon nigroviridis Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Actinopterygii; Neopterygii; Teleostei; Buteleostei; Neoteleostei; Acanthomozpha; Acanthopterygii; Percomorpha; Tetraodontiformes;	radontoidea; Tetraodontidae; Tetraodon. st Crollius, H., Jaillon, O., Dasilva, C., Bouneau, L., Fisher, C., not. A. Fizames, C. Wincker D. Brottier D. Onstier P.	and Weissenbach,J. of human gene number provided by genome raodon nigroviridis DNA sequence	Nat. Genet. 25 (2), 235-238 (2000) 20296633 10835645	<pre>st Crollius,H., Jaillon,O., Dasilva,C., Ozouf-Costaz,C., ammes,C., Fischer,C., Bonneau,L., Billault,A., Quetier,F.,</pre>	Characterization and repeat analysis of the compact genome of the freshwater pufferfish Tetraodon nigroviridis	2033983/ 10899143 3 (bases 1 to 1121) Genoscope.		- Web : www.genoscope.cns.ir) This sequence is a single read and was generated as part of a large scale clone-end sequencing project of the Tetraodon nigroviridis	p://www.genoscope.cns.fr/Tetraodon.   Location/Qualifiers   1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2	/organism="Tetraodon nigroviridis" /mol_type="genomic DNA" /db_xref="taxoon:99883" /clone="109N 9"	/clone_lib="G" /note="Genoscope sequence ID : COBGI09CGI0LP1~end : T7"	s: 46.7 Length: 1121 81.00 Matches: 33 Lty: 38.10% Conservative: 7
Db 444	Qy 52		Db 336 Qy 82	Db 279	RESULT 12 CNSO4H30 LOCUS DEFINITION T		ORGANISM T	T REFERENCE 1 AUTHORS R			AUTHORS R	TITLE CI JOURNAL G			COMMENT TI	hi FEATURES		ORIGIN	Alignment Scores: Pred. No.: Score: Percent Similarity:

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Vector: pCMV-SPORT6; Site 1: Not1; red); RNA source anonymous pool of 3 age 20 weeks, female age 24 weeks, Library is oligo-dT primed and (EcoRV site is destroyed upon sert size 1.7 kb, insert size range is normalized and enriched for a was constructed by C. Gruber ch Genetics tracking code 017. Note:
 mRNA linear EST 17-OCT-2001 ens cDNA clone IMAGE:5243769 5',
 SnThrLeuGluAsnGlyTyrPheLeuSer 33
 ||
|TITACITICIGITICCCAAAGITAAACA 978
 sgreregaaaagagccragaargaacreg 798
 CACGGCTTCAGCTTACCTGCTCAGGTGT 918
 1Cys-----GluAlaIleTrp 17
 uGluAspSerValThr-----Pro 53
 38
 iata; Vertebrata; Euteleostomi;
rrhini; Hominidae; Homo.
 ogies, Inc.
Technologies, Inc.
M.A.G.E. Consortium (LLNL)
istribution information can be
sortium/LLNL at:
 malian Gene Collection (MGC)
28
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Allogas, L. Chen, D. Chen, D. Chen, D., Shatshan, S., Tsegaye, G., Geer, K., Shetty, J., Shetty, J., Shatsman, S., Tsegaye, G., Geer, K., Shvartsbeyn, A., Gebregeorgis, E., Overton, L., Russell, D., Chen, D., Riggs, F., de Jong, P. and Fraser, C.M.
Rat BAC End Sequences from Library CHORI-230 EcoRI segment Unpublished (1999)

AL Unpublished (1999)

Other GSS: CH230-163F15.TV

Contact: Shaying Zhao

Contact: Shaying Zhao

Department of Eukaryotic Genomics

The Institute for Genomic Research

9712 Medical Center Dr., Rockville, MD 20850, USA

Tel: 301 838 0200

Email: Schaogetigr.org

Clones are derived from the rat BAC library CHORI-230

(http://www.chori.org/bacpac/rat230.thm). For BAC library

availability, please contact Pieter de Jong (pdejong@mail.cho.org).

Clones may be purchased from BACPAC Resources

(http://www.chori.org/bacpac/creting_information.htm). BAC end

page: http://www.tigr.org/tdb/bac_ends/rat/bac_end_intro.html

Plate: 163 row: F column: 15
 BH354326 CH230-163F15.TJ CHORI-230 Segment 1 Rattus norvegicus genomic clone CH230-163F15, genomic survey sequence.
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 34 ArgAsnLysGluAsnHisSerGlnProThrGlnSerSerLeuGluAspSerValThrPro 53
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 ----LeulleLeuGlyAla 79
 Bukaryota, Mětazoa, Chordata, Craniata, Vertebrata, Euteleostomi;
Mammalia, Butheria, Rodentia, Sciurognathi, Muridae, Murinae;
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 Rattus norvegicus (Norway rat)
Rattus norvegicus
 Location/Qualifiers
Pieter de Jong"
 BH354326
BH354326.1 GI:17285060
 Rattus.
1 (bases 1 to 504)
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79.00
46.38%
31.88%
17.10%
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Best Local Similarity:
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 GSS
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VERSION
 TITLE
JOURNAL
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 REFERENCE
 AUTHORS
 KEYWORDS
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Shaots, Shetty, J., Shatsman, S., Tsegaye, G., Geer, K.,
Shaots, Shetty, J., Gebregeorgis, E., Overton, L., Russell, D., Chen, D.,
Riggs, F., de Jong, P. and Fraser, C.M.
Rat BAC End Sequences from Library CHORI-230 EcoRI segment
Unpublished (1999)
Other GSS: CH230-163F16.TV
Contact: Shaying Zhao
Department of Eukaryotic Genomics
The Institute for Genomic Research
9712 Medical Center Dr., Rockville, MD 20850, USA
Tel: 301 838 0208
Email: szhao@tigr.org
Clones are derived from the rat BAC library CHORI-230
(http://www.chori.org/bacpac/rat230.htm). For BAC library
availability, please contact Pieter de Jong (pdejong@mail.cho.org).
Clones may be purchased from BACPAC Resources
// Letter A. // Letter C.
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 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae;
 39 HisserGlnProThrGlnSer-----SerLeuGluAspSerValThrProThr 54
 55 LysalavalLysThrThrGlyLysGlyIleValLysGlyArgAsnLeuAspSerArg---
 (http://www.chori.org/bacpac/or ering_information.htm). BAC er page: http://www.tigr.org/tdb/bac_ends/rat/bac_end_intro.html plate: 163 row: F column: 16 Seg primer: SP6 Class: BAC ends.
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 683
25
4
15
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47.17%
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 Percent Similarity:
 Rattus.
 Alignment Scores:
 GSS
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 RESULT 14
BH354332/c
 DEFINITION
 Pred. No.:
 ORGANISM
 TITLE
JOURNAL
COMMENT
 REFERENCE
AUTHORS
 ACCESSION
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 FEATURES
 VERSION
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EcoRI;
/cell type="Brain"
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_		

US-09-989-293A-377 (1-90) x BH354326 (1-504)

 $\Omega$ ga Zy

74 Gly-------ieulleLeuGlyAla 79

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214 GGGAGAAACTAATGCTTCCAACACAGACCCTCAGTGGAAAACTGCTTATGTGCAGGCAAA 155 

Search completed: December 2, 2004, 06:31:10 Job time: 2580 secs

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